Access DB# 5/15/7

SEARCH REQUEST FORM

Scientific and Technical Information Center

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Art Unit: /64 Phone Number 30 5-/89 Mail Box and Bldg/Room Location: 85-12 (041) Res	Serial Number: _9/55-1.9 77
Mail Box and Bldg/Room Location: 25 12 (04) Res	ults Format Preferred (circle) PAPER DISK E-MAIL
If m re than one search is submitted, please prioritia	ze searches in order of need.
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Please provide a detailed statement of the search topic, and describe Include the elected species or structures, keywords, synonyms, acror	
utility of the invention. Define any terms that may have a special m	eaning. Give examples or relevant citations, authors, etc, if
known. Please attach a copy of the cover sheet, pertinent claims, and	
Title of Invention: Compositions and Mothods	for generating and immune
Inventors (please provide full names):	tilizing alphavious -Basal voctor syste
	Dubensky. Mya Frolov. Jason Gardner
Earliest Priority Filing Date: April 14 1999	
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Date Searcher Picked Up: Bibliographic	Dr.Link
Date Completed: 09-19-01 Litigation	Lexis/Nexis
Searcher Prep & Review Time: 12 Fulltext	Sequence Systems
Clerical Prep Time: Patent Family	WWW/Internet
Online Time: Other	Other (specify)
PTO-1590 (1-2000)	

FILE 'CAPLUS' ENTERED AT 11:34:09 ON 19 SEP 2001

3621 SEA FILE=CAPLUS ABB=ON PLU=ON ALPHAVIRUS OR ALPHA

VIRUS OR VR2526 OR VR 2526 OR SINDBIS OR SEMLIKI FOREST

OR ROSS RIVER OR VENEZUEL? EQUINE (1W) VIRUS

37 SEA FILE=CAPLUS ABB=ON PLU=ON L1 AND (DENDRIT? OR

DC(S)DENDRIT?)

L6 ANSWER 1 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2001:647459 CAPLUS

TITLE:

L1

L6

Abundant GFP expression and LTP in hippocampal

acute slices by in vivo injection of

sindbis virus

AUTHOR (S):

D'Apuzzo, Massimo; Mandolesi, Georgia; Reis,

Gerald; Schuman, Erin M.

CORPORATE SOURCE:

California Institute of Technology, Pasadena,

CA, 91125, USA

SOURCE:

J. Neurophysiol. (2001), 86(2), 1037-1042

CODEN: JONEA4; ISSN: 0022-3077

PUBLISHER:

American Physiological Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Virus-mediated gene transfer into neurons is a powerful tool for the AB anal. of neuronal structure and function. Recombinant sindbis virus has been previously used to study protein function in hippocampal neuron cultures as well as in hippocampal organotypic slice cultures. Nevertheless, some concern still exists about the physiol. relevance of these cultured prepns. Acute hippocampal slices are a widely used prepn. for the study of synaptic transmission, but currently recombinant gene delivery is usually achieved only through time-consuming transgenic techniques. In this study, we show that a subregion of the CA1 area in acute hippocampal slices can be specifically altered to express a gene of interest. A sindbis virus vector carrying an enhanced green fluorescent protein (EGFP) reporter was injected in vivo into the hippocampus of adult rats. After 18 h, rats were killed, and acute hippocampal slices, infected in the CA1 field, were analyzed morphol. and electrophysiol. Infected slices showed healthy and stable electrophysiol. responses as well as long-term potentiation. In addn., infected pyramidal cells were readily recognized in living slices by two-photon imaging. Specifically, the introduction of an EGFP-Actin fusion protein greatly enhanced the detection of fine processes and dendritic spines. We propose this technique as an efficient tool for studying gene function in adult hippocampal neurons.

L6 ANSWER 2 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2001:636194 CAPLUS

TITLE:

Hybrid cell vaccines derived by fusion of an

allogeneic dendritic cells and a nondendritic cells and uses in tumor and infection therapy

infection therapy

INVENTOR(S):
PATENT ASSIGNEE(S):

Kanz, Lothar; Walden, Peter; Stuhler, Gernot Eberhard-Karls-Universitaet Tuebingen

Universitaetsklinikum, Germany

SOURCE:

PCT Int. Appl., 42 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

KIND DATE APPLICATION NO. DATE PATENT NO. **A**1 20010830 WO 2000-EP2433 20000320 WO 2001062902 AE, AG, AM, AU, AZ, BA, BB, BG, BR, BY, CA, CN, CR, CU, CZ, DM, DZ, EE, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, MA, MD, MG, MN, MW, MX, NO, NZ, PL, RU, SD, SG, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG EP 2000-105829 20000320 EP 1130088 20010905 **A**1 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO PRIORITY APPLN. INFO.: DE 2000-10009030 A 20000227 US 2000-185334 P 20000228

The present invention relates to methods and compns. for treating AB and preventing cancer and infectious disease using hybrid cells formed by fusion of allogeneic dendritic cells and autologous non-dendritic cells which shares at least one class I MHC (major histocompatibility complex) allele. Such hybrid cells combine the vigorous alloreactivity of mature dendritic cells with the specific antigenicity of autologous tumor cells, thereby eliciting a highly specific and vigorous cytotoxic T lymphocytes (CTL) response. The invention also provides the methods for making hybrid cell vaccines and evaluating its cytotoxicity. For rapid and large-scale generation of hybrids, electrofusion is established as a two-step procedure: in the first step, tumor cells and dendritic cells (DCs) were dielectrophoretically aligned to from cell-cell conjugates; in the second step, a fusion pulse was applied, yielding 10-15% hybrid cell formation. The invention demonstrates that vaccine with tumor celldendritic cell hybrid results in regression of human metastatic renal cell carcinoma.

REFERENCE COUNT:

6

REFERENCE(S):

(1) Celluzzi, C; JOURNAL OF IMMUNOLOGY 1998,

V160(7), P3081 CAPLUS

- (2) Dana Farber Cancer Inst Inc; WO 9846785 A 1998 CAPLUS
- (3) Guo, Y; SCIENCE 1994, V263(5146), P518 CAPLUS
- (4) Kugler, A; BRITISH JOURNAL OF UROLOGY 1998, V82(4), P487 MEDLINE
- (6) Kugler, A; NATURE MEDICINE 2000, V6(3), P332 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 3 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2001:563197 CAPLUS

TITLE:

Changes in calcium currents and GABAergic

spontaneous activity in cultured rat hippocampal neurons after a neurotropic influenza A virus

infection

AUTHOR (S):

Brask, J.; Owe-Larsson, B.; Hill, R. H.;

Kristensson, K.

CORPORATE SOURCE:

Department of Neuroscience, Karolinska

Institutet, Stockholm, Swed.

SOURCE:

Brain Res. Bull. (2001), 55(3), 421-429

CODEN: BRBUDU; ISSN: 0361-9230

PUBLISHER:

Elsevier Science Inc.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

In order to study mechanisms by which a neurotropic strain of influenza A virus (A/WSN/33) may affect neuronal function or cause nerve cell death, hippocampal cultures from embryonic rats were infected with this virus. Approx. 70% of the neurons in the infected cultures became immunopos. for viral antigens and showed reduced voltage-dependent Ca2+ currents in whole-cell patch clamp recordings, but no changes in other membrane properties or in cytosolic Ca2+ concn. were seen. These immunopos. neurons underwent apoptosis 3-4 days after infection. Ca2+ channel inhibitors had no significant effect on neuronal survival. The immunoneg. population of neurons survived, but displayed increased frequency of miniature inhibitory postsynaptic currents of .gamma.-amino-butyric acid origin compared with controls. The frequency of .alpha.-amino-hydroxy-5-methylisoxazole-4-propionic acid hydrobromide (AMPA) receptor-mediated miniature excitatory postsynaptic currents was not altered. Viral nucleoproteins, overexpressed using the Semliki Forest virus system, were localized to the dendritic spines as shown by double immunolabeling with actinin, but did not by themselves cause neuronal death or changes in synaptic transmission as measured by AMPA-mediated excitatory postsynaptic currents. Our results show that an influenza A virus infection can cause selective

neurophysiol. changes in hippocampal neurons and that these can persist even after the viral antigens have been cleared.

L6 ANSWER 4 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 20

2001:545880 CAPLUS

DOCUMENT NUMBER:

135:103358

TITLE:

Novel non-lab. virus strains with improved

oncolytic and/or gene delivery capabilities as

compared to lab. virus strains, and uses thereof

INVENTOR(S):

Coffin, Robert Stuart

PATENT ASSIGNEE(S):

Biovex Limited, UK PCT Int. Appl., 48 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

PA	PATENT NO.				KIND DATE													
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WO	2001	0535	06	A	2 :	2001	0726		W	0 20	01-G	B229		2001	0122			
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		CN,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,		
		GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,		
		LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,		
		PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,	TJ,	TM,	TR,	TT,	TZ,		
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		ТJ,	TM															
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		•						(GB 2	001-	430		A	2001	0106			
AB The invention provides novel non-lab. virus strains, esp. herpes																		

The invention provides novel non-lab. virus strains, esp. herpes viruses such as HSV, with improved oncolytic and/or gene delivery capabilities as compared to lab. virus strains. The inventors have shown that two clin. isolates of HSV1 (strains JS1 and BL1) have enhanced replication in some human tumor cell lines as compared to the lab. HSV1 strain 17+. Although the invention is exemplified using HSV, it is equally applicable for other viruses, such as adenovirus, picornavirus, retrovirus, or alphavirus. In the provided virus strains, the gene encoding ICP34.5 was deleted, and this resulted in enhanced growth in tumor cells as compared to a similarly engineered lab. strain. The viruses also are engineered to express human GM-CSF or some other immunomodulatory protein. The

invention also provides methods and compns. for the treatment of cancer.

L6 ANSWER 5 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 2001:354947 CAPLUS

DOCUMENT NUMBER: 135:120926

TITLE: Enhancement of Sindbis virus

self-replicating RNA vaccine potency by linkage of Mycobacterium tuberculosis heat shock protein

70 gene to an antigen gene

AUTHOR(S): Cheng, Wen-Fang; Hung, Chien-Fu; Chai, Chee-Yin;

Hsu, Keng-Fu; He, Liangmai; Rice, Charles M.;

Ling, Morris; Wu, T.-C.

CORPORATE SOURCE: Department of Pathology, Johns Hopkins Medical

Institutions, Baltimore, MD, 21205, USA

SOURCE: J. Immunol. (2001), 166(10), 6218-6226

CODEN: JOIMA3; ISSN: 0022-1767

PUBLISHER: American Association of Immunologists

DOCUMENT TYPE: Journal LANGUAGE: English

Recently, self-replicating RNA vaccines (RNA replicons) have emerged AB as an effective strategy for nucleic acid vaccine development. Unlike naked DNA vaccines, RNA replicons eventually cause lysis of transfected cells and therefore do not raise the concern of integration into the host genome. We evaluated the effect of linking human papillomavirus type 16 E7 as a model Ag to Mycobacterium tuberculosis heat shock protein 70 (HSP70) on the potency of Aq-specific immunity generated by a Sindbis virus self-replicating RNA vector, SINrep5. Our results indicated that this RNA replicon vaccine contg. an E7/HSP70 fusion gene generated significantly higher E7-specific T cell-mediated immune responses in vaccinated mice than did vaccines contg. the wild-type E7 gene. Furthermore, our in vitro studies demonstrated that E7 Ag from E7/HSP70 RNA replicon-transfected cells can be processed by bone marrow-derived dendritic cells and presented more efficiently through the MHC class I pathway than can wild-type E7 RNA replicon-transfected cells. More importantly, the fusion of HSP70 to E7 converted a less effective vaccine into one with significant potency against E7-expressing tumors. This antitumor effect was dependent on NK cells and CD8+ T cells. These results indicated that fusion of HSP70 to an Ag gene may greatly enhance the potency of self-replicating RNA vaccines.

REFERENCE COUNT:

41

REFERENCE(S):

- (1) Albert, M; J Exp Med 1998, V188, P1359 CAPLUS
- (2) Albert, M; Nature 1998, V392, P86 CAPLUS
- (4) Anthony, L; Vaccine 1999, V17, P373 CAPLUS
- (5) Basu, S; Int Immunol 2000, V12, P1539 CAPLUS

(6) Berglund, P; AIDS Res Hum Retrovir 1997, V13, P1487 CAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 6 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2001:205129 CAPLUS

DOCUMENT NUMBER:

134:365427

TITLE:

Infection of human dendritic cells by a sindbis virus replicon vector is

determined by a single amino acid substitution

in the E2-glycoprotein

AUTHOR(S):

Gardner, Jason P.; Frolov, Ilya; Perri, Silvia; Ji, Yaying; MacKichan, Mary Lee; zur Megede, Jan; Chen, Minchao; Belli, Barbara A.; Driver, David A.; Sherrill, Scott; Greer, Catherine E.; Otten, Gillis R.; Barnett, Susan W.; Liu, Margaret A.; Dubensky, Thomas W.; Polo, John M.

CORPORATE SOURCE:

Vaccines & Gene Therapy, Chiron Corporation,

Emeryville, CA, 94608, USA

SOURCE:

J. Virol. (2000), (74(24), 11849-11857

CODEN: JOVIAM; ISSN: 0022-538X

PUBLISHER:

American Society for Microbiology

DOCUMENT TYPE: LANGUAGE: Journal

English The ability to target antigen-presenting cells with vectors encoding AB desired antigens holds the promise of potent prophylactic and therapeutic vaccines for infectious diseases and cancer. this goal, we derived variants of the prototype alphavirus , Sindbis virus (SIN), with differential abilities to infect human dendritic cells. Cloning and sequencing of the SIN variant genomes revealed that the genetic determinant for human dendritic cell (DC) tropism mapped to a single amino acid substitution at residue 160 of the envelope glycoprotein E2. Packaging of SIN replicon vectors with the E2 glycoprotein from a DC-tropic variant conferred a similar ability to efficiently infect immature human DC, whereupon those DC were obsd. to undergo rapid activation and maturation. The SIN replicon particles infected skin-resident mouse DC in vivo, which subsequently migrated to the draining lymph nodes and upregulated cell surface expression of major histocompatibility complex and costimulatory mols. Furthermore, SIN replicon particles encoding human immunodeficiency virus type 1 p55Gag elicited robust Gag-specific T-cell responses in vitro and in vivo, demonstrating that infected DC maintained their ability to process and present replicon-encoded antigen. Interestingly, human and mouse DC were differentially infected by selected SIN variants, suggesting

differences in receptor expression between human and murine DC.

These data illustrate the tremendous potential of using a directed

approach in generating alphavirus vaccine vectors that target and activate antigen-presenting cells, resulting in robust antigen-specific immune responses.

REFERENCE COUNT:

50

REFERENCE(S):

- (1) Akbari, O; J Exp Med 1999, V189, P169 CAPLUS
- (2) Albert, M; Nature 1998, V392, P86 CAPLUS
- (3) Banchereau, J; Nature 1998, V392, P245 CAPLUS
- (4) Bender, A; J Immunol Methods 1996, V196, P121 CAPLUS
- (5) Bhardwaj, N; J Exp Med 1997, V186, P795 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 7 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2001:196378 CAPLUS

TITLE:

Enhancement of antitumor immune response in glioma models in mice by genetically modified

dendritic cells pulsed with
Semliki Forest virus-mediated

complementary DNA

AUTHOR(S):

Yamanaka, Ryuya; Zullo, Susan A.; Tanaka, Ryuichi; Blaese, Michael; Xanthopoulos,

Kleanthis G.

CORPORATE SOURCE:

Clinical Gene Therapy Branch, National Institutes of Health, Bethesda, MD, USA

SOURCE:

J_Neurosurg. (2001), 94(3), 474-481

CODEN: JONSAC; ISSN: 0022-3085

PUBLISHER:

American Association of Neurological Surgeons

DOCUMENT TYPE:

Journal

LANGUAGE:

English

The aim of this study was to further investigate dendritic cell (DC) -based immunotherapy for malignant glioma to improve its therapeutic efficacy. Dendritic cells were isolated from the bone marrow and pulsed with phosphate-buffered saline, tumor RNA, tumor lysate, Semliki Forest virus (SFV)-LacZ, SFV-mediated B16 complementary (c)DNA, or SFV-mediated 203 glioma cDNA, resp., to treat mice bearing tumors of the 203 glioma cell line. The results indicated that pre-immunization with DCs pulsed with the same type of cDNA as in the tumor by a self-replicating RNA vector (i.e., SFV) protected mice from tumor challenge, and that therapeutic immunization prolonged the survival of mice with established tumors. The SFV induced apoptosis in DCs and their death facilitated the uptake of apoptotic cells by other DCs, thus providing a potential mechanism for enhanced immunogenicity. Conclusions. Therapy with DCs that have been pulsed with SFV-mediated tumor cDNA may be an excellent

procedure for the development of new cancer vaccines.

23 REFERENCE COUNT: (1) Albert, M; Nature 1998, V392, P86 CAPLUS REFERENCE(S): (2) Ashley, D; J Exp Med 1997, V186, P1177 **CAPLUS** (3) Boczkowski, D; J Exp Med 1996, V184, P465 CAPLUS (4) Celluzzi, C; J Exp Med 1996, V183, P283 (5) Engelhardt, J; Hum Gene Ther 1993, V4, P759 **CAPLUS** ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 8 OF 37 CAPLUS COPYRIGHT 2001 ACS L6 2001:168165 CAPLUS ACCESSION NUMBER: 134:217978 DOCUMENT NUMBER: Antibody-dependent enhancement of TITLE: alphavirus vector transfection MacDonald, Gene H.; Johnston, Robert E. INVENTOR(S): University of North Carolina at Chapel Hill, USA PATENT ASSIGNEE(S): PCT Int. Appl., 66 pp. SOURCE: CODEN: PIXXD2 Patent DOCUMENT TYPE: English LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. DATE KIND DATE PATENT NO. 20010308 WO 2000-US23845 20000830 WO 2001016343 A1 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE,

The present invention provides compns. and methods for delivering a nucleotide sequence to a cell using an alphavirus vector that is complexed with an enhancing antibody that specifically binds to the alphavirus vector. Venezuelan Equine Encephalitis vectors are preferred. The cell may be a cell in vitro or in vivo. Alternatively, the cell may be removed from a subject, administered the alphavirus vector ex vivo and then administered to a

PRIORITY APPLN. INFO.:

Searcher: Shears 308-4994

BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

US 1999-151718

US 2000-177435

P 19990831

P 20000121

subject. Antigen-presenting cells are preferred, with dendritic cells being more preferred. Also provided are methods of producing an immune response in a subject, e.g., for producing an immune response against an antigen assocd. With a pathogen or for immunotherapy of cancer of tumors.

REFERENCE COUNT:

REFERENCE(S):

(1) Akzo Nobel Nv; EP 0659885 A 1995 CAPLUS

(2) Anon; http://wwwamb.casaccia.enea.it/glc/CD-table.htm 1995

(3) Linn, M; J GEN VIROL 1996, V77, P407 CAPLUS

(4) Univ North Carolina; WO 9532733 A 1995 CAPLUS

L6 ANSWER 9 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2001:135251 CAPLUS

DOCUMENT NUMBER:

134:294223

TITLE:

Enhancement of Sindbis virus

self-replicating RNA vaccine potency by targeting antigen to endosomal/lysosomal

compartments

AUTHOR(S):

Cheng, Wen-Fang; Hung, Chien-Fu; Hsu, Keng-Fu; Chai, Chee-Yin; He, Liangmei; Ling, Morris; Slater, Leigh A.; Roden, Richard B. S.; Wu,

T.-C.

CORPORATE SOURCE:

Department of Pathology, Johns Hopkins Medical

Institutions, Baltimore, MD, 21205, USA

SOURCE:

Hum. Gene Ther. (2001), 12(3), 235-252

CODEN: HGTHE3; ISSN: 1043-0342

PUBLISHER:

LANGUAGE:

Mary Ann Liebert, Inc.

DOCUMENT TYPE:

Journal English

Self-replicating RNA vaccines (RNA replicons) have emerged as an AB attractive approach for tumor immunotherapy. RNA replicons do not integrate into host chromosomes, eliminating the concern for oncogenicity assocd. with a DNA vaccine. In this study, the authors used human papillomavirus type 16 (HPV-16) E7 as a model antigen and evaluated E7-specific immunity generated by a Sindbis virus self-replicating RNA vector, SIN-rep5. Three different constructs were created to target E7 antigen to different cellular localizations: (1) E7, a cytosolic/nuclear protein; (2) Sig/E7, a secretory protein; (3) Sig/E7/LAMP-1, in which the authors linked the transmembrane and cytoplasmic regions of the lysosome-assocd. membrane protein 1 (LAMP-1) to E7 protein to target E7 to the endosomal/lysosomal compartment. The authors found that the RNA replicon vaccine contg. the Sig/E7/LAMP-1 fusion gene generated the highest E7-specific T cell-mediated immune responses and antitumor effects relative to RNA vaccines contg. either wild-type E7 or Sig/E7. Our in vitro studies demonstrated that E7 antigen from

Sig/E7/LAMP-1 RNA replicon-transfected apoptotic cells can be taken up by bone marrow-derived dendritic cells (DCs) and presented more efficiently through the MHC class I pathway than wild-type E7 RNA replicon-transfected apoptotic cells. Furthermore, the authors' data revealed that CD8+ T cells, CD4+ T cells, and NK cells were important for the antitumor effects generated by Sig/E7/LAMP-1 RNA vaccination. These results indicate that targeting antigen to the endosomal/lysosomal compartment via fusion to LAMP-1 may greatly enhance the potency of self-replicating RNA vaccines.

REFERENCE COUNT:

44

REFERENCE(S):

- (1) Albert, M; J Exp Med 1998, V188, P1359 CAPLUS
- (2) Albert, M; Nature 1998, V392, P86 CAPLUS
- (3) Berglund, P; AIDS Res Hum Retroviruses 1997, V13, P1487 CAPLUS
- (4) Berglund, P; Nat Biotechnol 1998, V16, P562 CAPLUS
- (5) Bigger, J; J Immunol 1998, V160, P5826 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 10 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:742266 CAPLUS

DOCUMENT NUMBER:

133:320989

TITLE:

Compositions and methods for generating an

immune response utilizing alphavirus

-based vector systems

INVENTOR(S):

Polo, John M.; Dubensky, Thomas W., Jr.; Frolov,

Ilya; Gardner, Jason P.; Otten, Gillis; Barnett,

Susan; Driver, David A.

PATENT ASSIGNEE(S):

Chiron Corporation, USA

SOURCE:

PCT Int. Appl., 83 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

1

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	CENT	NO.		KI	ND :	DATE			A.	PPLI	CATI	ON NC	o. :	DATE			
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WO	2000	0617	72	A:	2	20001019 WO 2000-US10722						22	20000414				
WO	2000	0617	72	A.	3	2001	0208										
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	
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		LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	
		RO.	RII.	SD.	SE.	SG.	ST.	SK.	SL.	TJ.	TM.	TR.	TT.	TZ.	UA.	UG.	

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US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF,
BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO.:
US 1999-129498 P 19990414
US 1999-148086 P 19990809
US 2000-191363 P 20000322

AB Methods are provided for generating immune responses utilizing
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alphavirus-based vector systems. Thus, Sindbis
virus is passaged 4 times in primary human dendritic cells
obtained from different donors, with intermediate plaque purifn. in
293 and BHK-21 cells. The resulting viral strains infect human
dendritic cells, primarily as a result of an amino acid
substitution at residue 160 of the E2 glycoprotein as compared to
the wild type. Other similar alphavirus-based systems,
such as Semliki Forest virus, Venezuelan
equine encephalitis virus, and Ross
Pivor virus, also may be readily substituted. Eukaryotic

River virus, also may be readily substituted. Eukaryotic layered vector initiation systems (ELVIS) are provided comprising a 5' promoter capable of initiating in vivo the 5' synthesis of alphavirus RNA from cDNA, a sequence which initiates transcription of alphavirus RNA following the 5'-promoter, a nucleic acid mol. which operably encodes all 4 alphaviral nonstructural proteins, an alphavirus RNA polymerase recognition sequence, and a 3' polyadenylate tract. The nonstructural proteins contain a mutation in at lease one of residues 346, 441, or 473 in nsP1; residues 438, 622, 634, or 715 in nsP2; residues 417, 456, or 505 in nsP3, and residues 266 in nsP4. The construction of a full-length cDNA clone, replicon vectors, and structural protein expression (packaging) cassettes from a human dendritic cell adapted alphavirus, such as

SinDCchiron virus, is readily accomplished. A wide variety of therapeutic proteins and antigens may be expressed from the alphavirus-based vector systems, as demonstrated using the HIV gag polypeptide as an example antigen.

L6 ANSWER 11 OF 37 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 2000:727180 CAPLUS

DOCUMENT NUMBER:

134:264711

TITLE:

Alphavirus DNA and particle replicons

for vaccines and gene therapy

AUTHOR (S):

Polo, J. M.; Gardner, J. P.; Ji, Y.; Belli, B. A.; Driver, D. A.; Sherrill, S.; Perri, S.; Liu,

M. A.; Dubensky, T. W., Jr.

CORPORATE SOURCE:

SOURCE:

Vaccines & Gene Therapy, Emeryville, CA, USA
Dev. Biol. (2000), 104 (Development and Clinical

Progress of DNA Vaccines), 181-185

CODEN: DBEIAI; ISSN: 1424-6074

PUBLISHER:

S. Karger AG

DOCUMENT TYPE:

Journal; General Review

LANGUAGE:

English

equine encephalitis virus (VEE), are currently

AB A review with 15 refs. Alphaviruses have several features that make them attractive as gene delivery platforms, and vectors derived principally from Sindbis virus (SIN), Semliki Forest virus (SFV), and Venezuelan

being developed as prophylactic and therapeutic vaccines for infectious diseases and cancer. Alphavirus vectors,

termed "replicons", retain the nonstructural protein genes encoding the viral replicase, that in turn program high level cytoplasmic amplification of the vector RNA. We have developed plasmid DNA and recombinant vector particle delivery systems derived from the prototype alphavirus, SIN. Each system uses RNA

prototype alphavirus, SIN. Each system uses RNA polymerase II-based expression of alphavirus genome components and both vector formats are highly efficacious towards inducing robust antigen-specific immune responses in vaccinated animals. To increase the potency of SIN vector particles, which are not known to be lymphotropic, the tropism was re-directed for efficient infection of dendritic cells, both in vitro and in vivo.

REFERENCE COUNT:

15

REFERENCE(S):

- (1) Berglund, P; Nature Biotech 1998, V16, P562 CAPLUS
- (2) Cella, M; J Exp Med 1999, V189, P821 CAPLUS
- (3) Dubensky, T; J Virol 1996, V70, P508 CAPLUS
- (4) Frolov, I; J Virol 1994, V68, P1721 CAPLUS
- (5) Frolov, I; J Virol 1997, V71, P2819 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 12 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:457216 CAPLUS

DOCUMENT NUMBER:

133:100437

TITLE:

Polynucleotides encoding antigenic HIV type C

Gag- and/or Env-containing polypeptides for AIDS

vaccine development

INVENTOR(S):

Barnett, Susan; Zur, Megede Jan

PATENT ASSIGNEE(S):

Chiron Corp., USA

SOURCE:

PCT Int. Appl., 113 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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20000706
                                           WO 1999-US31273 19991230
     WO 2000039304
                       A2
     WO 2000039304
                       A3
                            20010118
             AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU,
             CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL,
             IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV,
             MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG,
             SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZA, ZW, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                        US 1998-114495
                                                         P 19981231
PRIORITY APPLN. INFO.:
                                        US 1999-152195
                                                         P 19990901
     The present invention relates to polynucleotides encoding
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The present invention relates to polynucleotides encoding immunogenic HIV type C Gag- and/ofr Env-contg. polypeptides which can be used as AIDS vaccines. The coding sequences of HIV-1 Env, Gag, Gag-protease and Gag-polymerase, and some inhibitory (or instability) elements (INS) located within these coding sequences are modified to construct more efficient expression vectors. Methods of generation of packaging cell lines, and prodn. of Gag-and/or Env-contg. proteins, analyzing the synthetic gene expression, immunizing animals with the Sindbis virus expression constructs, and evaluating their immunogenicity are described.

L6 ANSWER 13 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 2000:431421 CAPLUS

TITLE: Langerhans cells migrate to local lymph nodes

following cutaneous infection with an arbovirus

AUTHOR(S): Johnston, Linda J.; Halliday, Gary M.; King,

Nicholas J. C.

CORPORATE SOURCE: Departments of Pathology, University of Sydney,

Sydney, 2006, Australia

SOURCE: J. Invest. Dermatol. (2000), 114(3), 560-568

CODEN: JIDEAE; ISSN: 0022-202X

PUBLISHER: Blackwell Science, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

Whereas there has been recent interest in interactions between dendritic cells and pathogenic viruses, the role of dendritic cells in the initiation of protective immunity to such organisms has not been elucidated. The aim of this study was to examine whether a resident dendritic cell population in the skin, Langerhans cells, respond to cutaneous viral infections which are effectively cleared by the immune system. We therefore characterized the ability of Langerhans cells to migrate to local draining lymph nodes following infection with the arthropod-borne viruses, West Nile virus or Semliki Forest virus. The data show that major histocompatibility complex class

II+/NLDC145+/E-cadherin+ Langerhans cell nos. are increased in the draining lymph nodes of infected mice and this increase is accompanied by a concomitant decrease in the Langerhans cell d. in the epidermis. Langerhans cell migration is assocd. with an accumulation of leukocytes in the lymph node, which is one of the earliest events in the initiation of an immune response. Both the migratory response and the draining lymph node leukocyte accumulation were abrogated if UV-inactivated instead of live viruses were used, suggesting the activation and subsequent migration of Langerhans cells requires a live, replicating antigen. Our findings are likely to have wider implications for the development of epidermally delivered vaccines and suggest that mobilization of dendritic cells may be involved in the development of immune responses to arthropod-borne viruses.

REFERENCE COUNT:

51

REFERENCE(S):

- (1) Aiba, S; J Immunol 1990, V145, P2791 CAPLUS
- (2) Aiba, S; J Invest Dermatol 1993, V100, P143 CAPLUS
- (3) Becker, Y; Virus Genes 1995, V9, P133 CAPLUS
- (4) Blacklaws, B; J Virol 1995, V69, P1400 CAPLUS
- (5) Borkowski, T; Eur J Immunol 1994, V24, P2767 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 14 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:402017 CAPLUS

DOCUMENT NUMBER:

133:54574

TITLE:

Recombinant vectors expressing multiple

costimulatory molecules, host cell infection,

and uses in immunogenic applications

INVENTOR (S):

Schlom, Jeffrey; Hodge, James; Panicali, Dennis

PATENT ASSIGNEE(S): United States Dept. of Health and Human

Services, USA; Therion Biologics Corporation

SOURCE: PCT Int. Appl., 188 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.		KINI	D I	DATE			Al	PLIC	CATIO	ON NO). 1	DATE		
								- 						
WO 20000344	94	A1	2	20000	615		W	199	99-US	52686	56	1999:	1112	
W: AE,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,
CU,	CZ,	DE, 1	DK,	DM,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,
ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,
LU,	LV,	MA, I	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,

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SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ,
             VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                        US 1998-111582
                                                         P 19981209
PRIORITY APPLN. INFO.:
     The present invention provides recombinant vectors encoding and
AB
     expressing at least three or more costimulatory mols and host cells
     infected by the vector. The recombinant vector may addnl. contain a
     gene encoding one or more target antigens or immunol. epitope as
     well as cytokine, chemokine, or Flt-3L. A method of making a
     recombinant poxvirus, of enhancing an immune response of an
     individual by administering a recombinant vector, and of treating or
     preventing a disease by activating a T lymphocyte, are also
     presented. Further describes are a method of making a progenitor
     dendritic cell or dendritic cell, of assesing the
     efficacy of a vaccine against a target antigen, and of screening for
     novel immunogenic peptides. The synergistic effect of these
     costimulatory mols. on the enhanced activation of T cells was
     demonstrated. The degree of T-cell activation using recombinant
     vectors contg. genes encoding three costimulatory mols. was far
     greater than the sum of recombinant vector constructs contg. one
     costimulatory mol. and greater than the use of two costimulatory
     mols. Results employing the triple costimulatory vectors were most
     dramatic under conditions of either low levels of first signal or
     low stimulator to T-cell ratios. This phenomenon was obsd. with
     both isolated CD4+ and CD8+ T cells. The recombinant vectors of the
     present invention are useful as immunogenes and vaccines against
     cancer and pathogenic micro-organisms, and in providing host cells,
     including dendritic cells and splenocytes with enhanced
     antigen-presenting functions.
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REFERENCE COUNT:

4

REFERENCE(S):

- (1) Hodge, J; Cancer Res 1999, V59, P5800 CAPLUS
- (2) Keting, C; US 5738852 A 1998 CAPLUS
- (3) Therion Biolog Corp; WO 9804727 A 1998 CAPLUS
- (4) US Health; WO 9610419 A 1996 CAPLUS

L6 ANSWER 15 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:254039 CAPLUS

DOCUMENT NUMBER:

132:289590

INVENTOR(S):

Peptide-enhanced cationic lipid transfections Hawley-Nelson, Pamela; Lan, Jianqing; Shih, Pojen; Jessee, Joel A.; Schifferli, Kevin P.;

Gebeyehu, Gulilat

PATENT ASSIGNEE(S):

Life Technologies, Inc., USA

SOURCE:

TITLE:

U.S., 103 pp., Cont.-in-part of U.S. 5,736,392.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 4

PATENT INFORMATION:

PA'	TENT	NO.		KI	ND	DATE			A	PPLI	CATI	ON NO	ο.	DATE		
	- - - -								-	- -						
US	6051	429		Α		2000	0418		U	S 19	97-8	1820	0	1997	0314	
US	5736	392		A		1998	0407	,	U	S 19	96-6	5813	0	1996	0604	
WO	9840	502		A	1	1998	0917		W	0 19	98-U	S523	2	1998	0316	
	W:	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,
		DE,	DK,	EE,	ES,	FI,	GB,	GE,	GH,	GM,	GW,	HU,	ID,	IL,	IS,	JP,
		KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,	LT,	ĻU,	LV,	MD,	MG,	MK,
		MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK,	SL,
		ТJ,	TM,	TR,	TT,	UA,	UG,	UZ,	VN,	YU,	ZW,	AM,	AZ,	BY,	KG,	KZ,
		MD,	RU,	ТJ,	TM											
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SZ,	ŪG,	ZW,	AT,	BE,	CH,	DE,	DK,	ES,
		FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,	CG,
-		CI,	CM,	GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG					
AU	9865	622		A	1	1998	0929		A	U 19	98-6	5622		1998	0316	
EP	1007	699		A	1	2000	0614		E	P 19	98-9	1173	7	1998	0316	
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,
		PT,	ΙE,	FI												
PRIORIT	Y APP	LN.	INFO	. :					US 1	995-	4773	54	B2	1995	0607	
			•						US 1	996-	6581	3 0	A2	1996	0604	
									US 1	997-	8182	00	Α	1997	0314	
									WO 1	998-1	ÚS52	32	W	1998	0316	
AB The	e pre	sent	inve	enti	on p	rovi	des (comp	ns.	usef	ul f	or t	rans	fect	ing	

AB The present invention provides compns. useful for transfecting eukaryotic cells comprising nucleic acid complexes with peptides, wherein the peptide is optionally covalently coupled to a nucleic acid-binding group, and cationic lipids or dendrimers as transfection agents. The invention also provides transfection compns. in which a peptide is covalently linked to the transfection agent (lipid, cationic lipid or dendrimer). Inclusion of peptides or modified-peptides in transfection compns. or covalent attachment of peptides to transfection agents results in enhanced transfection efficiency. Methods for the prepn. of transfection compns. and methods of using these transfection compns. as intracellular delivery agents and extracellular targeting agents are also disclosed.

REFERENCE COUNT:

46

REFERENCE(S):

- (2) Anon; EP 0359347 1990 CAPLUS
- (4) Anon; EP 0544292 1992 CAPLUS
- (5) Anon; WO 9213570 1992 CAPLUS
- (7) Anon; WO 9307282 1993 CAPLUS
- (8) Anon; WO 9307283 1993 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 16 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 2000:183080 CAPLUS

DOCUMENT NUMBER: 132:320767

TITLE: Alpha/Beta interferon protects adult mice from

fatal Sindbis virus infection and is

an important determinant of cell and tissue

tropism

AUTHOR(S): Ryman, Kate D.; Klimstra, William B.; Nguyen,

Khuong B.; Biron, Christine A.; Johnston, Robert

E.

CORPORATE SOURCE: Department of Microbiology and Immunology,

University of North Carolina at Chapel Hill,

Chapel Hill, NC, 27599, USA

SOURCE: J. Virol. (2000), 74(7), 3366-3378

CODEN: JOVIAM; ISSN: 0022-538X

PUBLISHER: American Society for Microbiology

DOCUMENT TYPE: Journal LANGUAGE: English

Infection of adult 129 Sv/Ev mice with consensus Sindbis AB virus strain TR339 is subclin. due to an inherent restriction in early virus replication and viremic dissemination. By comparing the pathogenesis of TR339 in 129 Sv/Ev mice and .alpha./.beta. interferon receptor null (IFN-.alpha./.beta.R-/-) mice, the authors have assessed the contribution of IFN-.alpha./.beta. in restricting virus replication and spread and in detg. cell and tissue tropism. In adult 129 Sv/Ev mice, s.c. inoculation with 100 PFU of TR339 led to extremely low-level virus replication and viremia, with clearance under way by 96 h postinoculation (p.i.). In striking contrast, adult IFN-.alpha./.beta.R-/- mice inoculated s.c. with 100 PFU of TR339 succumbed to the infection within 84 h. By 24 h p.i. a high-titer serum viremia had seeded infectious virus systemically, coincident with the systemic induction of the proinflammatory cytokines interleukin-12 (IL-12) p40, IFN-.gamma., tumor necrosis factor .alpha., and IL-6. Replicating virus was located in macrophage-dendritic cell (DC)-like cells at 24 h p.i. in the draining lymph node and in the splenic marginal zone. By 72 h p.i. virus replication was widespread in macrophage-DC-like cells in the spleen, liver, lung, thymus, and kidney and in fibroblast-connective tissue and periosteum, with sporadic IFN-.alpha./.beta.-mediated restriction of TR339 neuroinvasion. infection was mimicked in vitro in peritoneal exudate cells from 129 Sv/Ev vs. IFN-.alpha./.beta.R-/- mice. Thus, IFN-.alpha./.beta. protects the normal adult host from viral infection by rapidly conferring an antiviral state on otherwise permissive cell types, both locally and systemically. Ablation of the IFN-.alpha./.beta. system alters the apparent cell and tissue tropism of the virus and renders macrophage-DC-lineage cells permissive to infection.

REFERENCE COUNT:

65

REFERENCE(S):

- (3) Biron, C; Curr Opin Microbiol 1999, V2, P374 CAPLUS
- (5) Cella, M; J Exp Med 1999, V189, P821 CAPLUS
- (6) Charles, P; Virology 1995, V208, P662 CAPLUS
- (7) Ciavarra, R; J Immunol 1997, V158, P1749 **CAPLUS**
- (8) Cousens, L; J Exp Med 1999, V189, P1315 **CAPLUS**

ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 17 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:141420 CAPLUS

TITLE:

Papers to Appear in Forthcoming Issues

AUTHOR (S):

Anon.

SOURCE:

Cell. Immunol. (2000), 199(2), 138

CODEN: CLIMB8; ISSN: 0008-8749

PUBLISHER:

Academic Press

DOCUMENT TYPE:

Journal; Miscellaneous

English LANGUAGE: (Received and Accepted Dates Follow Title) Regulation of CS1 AB Fibronectin Expression and Function by IL-1 in Endothelial Cells. David L. Boyle, Yan Shi, Steffen Gay, and Gary S. Firestein. (Received 7/23/99; accepted 1/5/00.) The Autoimmune Accelerating Yaa Mutation Does Not Accelerate Murine AIDS. Ambros W. Hugin, Liliane Fossati-Jimack, and Shozo Izui. (Received 8/18/99; accepted 1/5/00.) Lectin Ligands on Human Dendritic Cells and Identification of a Peanut Agglutinin Pos. Subset in Blood. Sherbini, B. Hock, D. Fearnley, A. McLellan, S. Vuckovic, and D. N. J. Hart. (Received 9/9/99; accepted 1/5/00.) B Cells and Antibodies in the Pathogenesis of Myelin Injury in Semliki Forest Virus Encephalomyelitis. Tamar A. Smith-Norowitz, Raymond A. Sobel, and Foroozan Mokhtarian. (Received 9/24/99; accepted 1/5/00.) The CD40-Inducible Bcl-2 Family Member A1 Protects B Cells from Antigen Receptor-Mediated Apoptosis. Andrew Craxton, Peter I. Chuang, Geraldine Shu, John M. Harlan, and Edward A. Clark. (Received 12/13/99; accepted 1/9/00.) In Vitro Characterization of Five Humanized OKT3 Effector Function Variant Antibodies. Danlin Xu, Maria-Luisa Alegre, Sally S. Varga, Annette L. Rothermel, Alexander M. Collins, Virginia L. Pulito, Lewis S. Hanna, Kevin P. Dolan, Paul W. H. I. Parren, Jeffrey A. Bluestone, Linda K. Jolliffe, and Robert A. Zivin. (Received 8/12/99; accepted 1/10/00.)Differential Control of Neonatal Tolerance by Antigen Dose vs. Extended Exposure and Adjuvant. Booki Min, Kevin L. Legge, Jacque C. Caprio, Lequn Li, Randal Gregg, and Habib Zaghouani. (Received 9/29/99; accepted 1/10/00.). (c) 2000 Academic Press.

ANSWER 18 OF 37 CAPLUS COPYRIGHT 2001 ACS

DOCUMENT NUMBER:

132:263829

TITLE:

Stimulation of cytotoxic T cells against

idiotype immunoglobulin of malignant lymphoma with protein-pulsed or idiotype-transduced

dendritic cells

AUTHOR (S):

Osterroth, Frank; Garbe, Annette; Fisch, Paul;

Veelken, Hendrik

CORPORATE SOURCE:

Departments of Hematology/Oncology and

Pathology, Freiburg University Medical Center,

Freiburg, D-79106, Germany

SOURCE:

Blood (2000), 95(4), 1342-1349

CODEN: BLOOAW; ISSN: 0006-4971

PUBLISHER:

American Society of Hematology

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Because of their hypervariable regions and somatic mutations, the AB antigen receptor mols. of lymphomas (idiotypes) are tumor-specific antigens and attractive targets for anti-lymphoma immunotherapy. For the optimal induction of human idiotype-specific cytotoxic T cells (CTL), idiotype was presented to CD8+ peripheral blood mononuclear cells by monocyte-derived autologous dendritic cells (DC) after the endocytosis of idiotype protein or by idiotype-expressing DC. Recombinant idiotype was obtained as a functionally folded Fab fragment by periplasmic expression in Escherichia coli. Idiotype-expressing DC were generated by transduction with recombinant Semliki forest virus vectors encompassing heavy- or light-chain idiotype genes. Autologous lymphoblastoid cell lines stably transfected with Epstein-Barr virus-based idiotype expression vectors were used as target cells to detect idiotype-specific lysis. CTL stimulated with idiotype-loaded DC showed strong specific, CD8-mediated, and major histocompatibility complex (MHC) class I-restricted cytotoxicity against autologous heavy- and light-chain idiotype. stimulation with idiotype-transduced DC resulted in only moderate natural killer cell activity. These data confirm the existence of idiotype-specific CTL in patients with lymphoma, define a "good manufg. practice" -compatible protocol for the generation of these cells without the requirement of viable lymphoma cells, and favor the processing of exogenous antigen over DC transduction for the induction of MHC I-restricted CTL against idiotypes with unknown antigenicity.

REFERENCE COUNT:

43

REFERENCE(S):

- (1) Banchereau, J; Nature 1998, V392, P245 CAPLUS
- (3) Boon, T; J Exp Med 1996, V183, P725 CAPLUS
- (5) Cella, M; Curr Opin Immunol 1997, V9, P10 CAPLUS
- (7) Fernandez, N; Nat Med 1999, V5, P405 CAPLUS

Searcher :

Shears

308-4994

(8) Fields, P; J Mol Med 1996, V74, P673 CAPLUS ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 19 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

CORPORATE SOURCE:

2000:42478 CAPLUS

TITLE:

Role of dendritic cell targeting in

venezuelan equine encephalitis

virus pathogenesis

AUTHOR(S):

MacDonald, Gene H.; Johnston, Robert E. Department of Microbiology and Immunology, University of North Carolina at Chapel Hill

School of Medicine, Chapel Hill, NC, 27599-7290,

IISA

SOURCE:

J. Virol. (2000), 74(2), 914-922 CODEN: JOVIAM; ISSN: 0022-538X American Society for Microbiology

PUBLISHER: DOCUMENT TYPE:

Journal English

LANGUAGE:

AB

The initial steps of Venezuelan equine encephalitis virus (VEE) spread from inoculation in the skin to the draining lymph node have been characterized. By using green fluorescent protein and immunocytochem., dendritic cells in the draining lymph node were detd. to be the primary target of VEE infection in the first 48 h following inoculation. VEE viral replicon particles, which can undergo only one round of infection, identified Langerhans cells to be the initial set of cells infected by VEE directly following inoculation. These cells are resident dendritic cells in the skin, which migrate to the draining lymph node following activation. A point mutation in the E2 glycoprotein gene of VEE that renders the virus avirulent and compromises its ability to spread beyond the draining lymph blocked the appearance of virally infected dendritic cells in the lymph node in vivo. A second-site suppressor mutation that restores viral spread to lymphoid tissues and partially restore virulence likewise restored the ability of VEE to infect dendritic cells in vivo.

REFERENCE COUNT:

35

REFERENCE(S):

- (1) Banchereau, J; Nature 1998, V392, P245 CAPLUS
- (2) Bhardwaj, N; J Exp Med 1997, V186, P795 CAPLUS
- (3) Borrow, P; J Virol 1995, V69, P1059 CAPLUS
- (4) Caley, I; J Virol 1997, V71, P3031 CAPLUS
- (5) Charles, P; Virology 1995, V208, P662 CAPLUS

308-4994

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 20 OF 37 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 2000:29999 CAPLUS

Searcher : Shears

DOCUMENT NUMBER: 132:333014

TITLE: DNA and RNA-based vaccines: principles, progress

and prospects

AUTHOR(S): Leitner, Wolfgang W.; Ying, Han; Restifo,

Nicholas P.

CORPORATE SOURCE: National Cancer Institute, National Institutes

of Health, Bethesda, MD, 20892-1502, USA

SOURCE: Vaccine (1999), 18(9-10), 765-777

CODEN: VACCDE; ISSN: 0264-410X

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review with 42 refs. DNA vaccines were introduced less than a ABdecade ago but have already been applied to a wide range of infectious and malignant diseases. Here we review the current understanding of the mechanisms underlying the activities of these new vaccines. We focus on recent strategies designed to enhance their function including the use of immunostimulatory (CpG) sequences, dendritic cells (DC), co-stimulatory mols. and cytokine- and chemokine-adjuvants. Although genetic vaccines have been significantly improved, they may not be sufficiently immunogenic for the therapeutic vaccination of patients with infectious diseases or cancer in clin. trials. One promising approach aimed at dramatically increasing the immunogenicity of genetic vaccines involves making them "self-replicating". This can be accomplished by using a gene encoding RNA replicase, a polyprotein derived from alphaviruses, such as Sindbis virus. Replicase-contg. RNA vectors are significantly more immunogenic than conventional plasmids, immunizing mice at doses as low as 0.1 .mu.g of nucleic acid injected once i.m. Cells transfected with "self-replicating" vectors briefly produce large amts. of antigen before undergoing apoptotic death. This death is a likely result of requisite double-stranded (ds) RNA intermediates, which also have been shown to super-active DC. Thus, the enhanced immunogenicity of "self-replicating" genetic vaccines may be a result of the prodn. of pro-inflammatory dsRNA, which mimics an RNA-virus infection of host cells.

REFERENCE COUNT:

142

REFERENCE(S):

- (1) Albert, M; Nature 1998, V392, P86 CAPLUS
- (2) Barry, M; Vaccine 1997, V15, P788 CAPLUS
- (3) Bell, A; Virology 1997, V232, P241 CAPLUS
- (4) Berglund, P; Nat Biotech 1998, V16, P562 CAPLUS
- (5) Berglund, P; Trends Biotechnol 1996, V14, P130 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 21 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1999:659264 CAPLUS

DOCUMENT NUMBER: 131:285394

TITLE: Methods and modified cells for the treatment of

cancer

INVENTOR(S): MacDonald, Gene H.; Martin, Brian K.; Johnston,

Robert E.; Ting, Jenny P-y

PATENT ASSIGNEE(S): University of North Carolina at Chapel Hill,

USA; Ting, Jenny P.-Y.

SOURCE: PCT Int. Appl., 46 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	rent 1	NO.		KI	ND :	DATE			A -	PPLI	CATI	ON NO	o. :	DATE		
WO	9951	263		A:	2	1999:	1014		W	0 19	99-U	S770	4	1999	0408	
WO	9951	263		A.	3	2000	0203									
	W:	AL,	AM,	AT,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,
		CZ,	CZ,	DE,	DE,	DK,	DK,	EE,	EE,	ES,	FI,	FI,	GB,	GE,	GH,	GM,
		HR,	HU,	ID,	IL,	IN,	IS,	JP,	ĶΕ,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,
		LS,	LT,	LU,	LV,	MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,
		RU,	SD,	SE,	SG,	SI,	SK,	SK,	SL,	ТJ,	TM,	TR,	TT,	UA,	UG,	US,
		UΖ,	VN,	YU,	ZW,	AM,	AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM		
	RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,
		DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,
		CF,	CG,	CI,	CM,	GA,	GN,	GW,	ML,	MR,	NE,	SN,	TD,	TG		
AU	9937	440		A.	1	1999	1025		A	U 19	99-3	7440		1999	0408	
EP	1069	908		A	2	2001	0124		E	P 19	99-9	1980	3	1999	0408	
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,
		PT,	IE,	FI					•							
PRIORITY	Y APP	LN.	INFO	. :					US 1	998-	8109	2	P	1998	0408	
									WO 1	999-	US77	04	W	1999	0408	

The present invention provides methods of preventing and/or treating cancers (including tumors). In one preferred embodiment, the invention is practiced to induce regression of an existing cancer or tumor and/or to prevent metastasis and/or to prevent growth of metastatic nodules. In other preferred embodiments, the invention may be used as a prophylaxis to prevent the development of primary cancers through a childhood or adult vaccination program against specific tumor antigens for cancers with high incidences. In an alternate preferred embodiment, the present invention provides methods of establishing an immune response against a universal artificial tumor antigen through a childhood or adult vaccine program, thus providing a long-term immune response that can be utilized at any point to treat any cancer which develops later in

life. The present invention also provides cancer and tumor cells stably expressing an artificial antigen, preferably an artificial cell-surface antigen.

L6 ANSWER 22 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1999:461772 CAPLUS

DOCUMENT NUMBER:

131:209757

TITLE:

Recombinant Semliki Forest

virus and Sindbis virus efficiently

infect neurons in hippocampal slice cultures Ehrengruber, Markus U.; Lundstrom, Kenneth; Schweitzer, Christophe; Heuss, Christian;

Schlesinger, Sondra; Gahwiler, Beat H.

CORPORATE SOURCE:

Brain Research Institute, University of Zurich,

Zurich, CH-8057, Switz.

SOURCE:

AUTHOR(S):

Proc. Natl. Acad. Sci. U. S. A. (1999), 96(12),

7041-7046

CODEN: PNASA6; ISSN: 0027-8424

PUBLISHER:

National Academy of Sciences

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Gene transfer into nervous tissue is a powerful tool for the anal. AB of gene function. By using a rat hippocampal slice culture prepn., we show here that Semliki Forest virus (SFV) and Sindbis virus (SIN) vectors are useful for the effective infection of neurons. The stratum pyramidale and/or the granular cell layer were injected with recombinant virus encoding .beta.-galactosidase (LacZ) or green fluorescent protein (GFP). By using low concns. of injected SFV-LacZ or SIN-LacZ, we detected LacZ staining of pyramidal cells, interneurons, and granule cells. About 60% of the infected cells showed clear neuronal morphol.; thus, relatively few glial cells expressed the transgene. Expression of GFP from SFV and SIN vectors gave similar results, with an even higher percentage (>90%) of the GFP-pos. cells identified as neurons. Infected pyramidal cells were readily recognized in living slices, displaying GFP fluorescence in dendrites of up to fourth order and in dendritic spines. They appeared morphol. normal and viable at 1-5 days postinfection. We conclude that both SFV and SIN vectors efficiently transfer genes into neurons in hippocampal slice cultures. In combination with the GFP reporter, SFV and SIN vectors will allow the physiol. examn. of identified neurons that have been modified by overexpression or suppression of a specific gene product.

REFERENCE COUNT:

38

REFERENCE(S):

- (1) Ankarcrona, M; Neuron 1995, V15, P961 CAPLUS
- (3) Berglund, P; Biotechnology 1993, V11, P916 CAPLUS
- (4) Bredenbeek, P; J Virol 1993, V67, P6439

CAPLUS

- (6) Davis, N; Proc Natl Acad Sci USA 1986, V83, P6771 CAPLUS
- (8) Ehrengruber, M; Methods Enzymol 1998, V293, P483 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 23 OF 37 CAPLUS COPYRIGHT 2001 ACS L6

ACCESSION NUMBER:

1999:444820 CAPLUS

DOCUMENT NUMBER:

131:198315

TITLE:

Cancer therapy using a self-replicating RNA

vaccine

AUTHOR(S):

Ying, Han; Zaks, Tal Z.; Wang, Rong-Fu; Irvine, Kari R.; Kammula, Udai S.; Marincola, Francesco M.; Leitner, Wolfgang W.; Restifo, Nicholas P.

CORPORATE SOURCE:

Surgery Branch, National Cancer Institute,

Bethesda, MD, 20892-1502, USA

SOURCE:

Nat. Med. (N. Y.) (1999), 5(7), 823-827

CODEN: NAMEFI; ISSN: 1078-8956

PUBLISHER:

Nature America

DOCUMENT TYPE:

Journal

English LANGUAGE: 'Naked' nucleic acid vaccines are potentially useful candidates for AB the treatment of patients with cancer, but their clin. efficacy has vet to be demonstrated. We sought to enhance the immunogenicity of a nucleic acid vaccine by making it 'self-replicating'. We accomplished this by using a gene encoding an RNA replicase polyprotein derived from the Semliki forest virus, in combination with a model antigen. A single i.m. injection of a self-replicating RNA immunogen elicited antigen-specific antibody and CD8+ T-cell responses at doses as low as 0.1 .mu.g. Pre-immunization with a self-replicating RNA vector protected mice from tumor challenge, and therapeutic immunization prolonged the survival of mice with established tumors. The self-replicating RNA vectors did not mediate the prodn. of substantially more model antigen than a conventional DNA vaccine did in vitro. However, the enhanced efficacy in vivo correlated with a caspase-dependent apoptotic death in transfected cells. This death facilitated the uptake of apoptotic cells by dendritic cells, providing a potential mechanism for enhanced immunogenicity. Naked, non-infectious, self-replicating RNA may be an excellent candidate for the development of new cancer vaccines.

REFERENCE COUNT:

23

REFERENCE(S):

- (1) Albert, M; Nature 1998, V392, P86 CAPLUS
- (2) Atkins, G; Mol Biotechnol 1996, V5, P33 CAPLUS
- (3) Berglund, P; Nature Biotechnol 1998, V16, P562 CAPLUS

Shears 308-4994 Searcher :

(4) Cella, M; J Exp Med 1999, V189, P821 CAPLUS

(5) Chappell, D; Cancer Res 1999, V59, P59 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 24 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1999:401694 CAPLUS

DOCUMENT NUMBER:

131:43583

TITLE:

Venezuelan equine

encephalitis **virus** vectors expressing tumor-associated antigens to induce cancer

immunity

INVENTOR(S):

Hippenmeyer, Paul J.

PATENT ASSIGNEE(S):

G.D. Searle & Co., USA

SOURCE:

PCT Int. Appl., 40 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO	DATE
WO 9930734	λ1 19990624	WO 1998-US2572	5 19981214
W: AL, AM,	AT, AU, AZ, BA,	BB, BG, BR, BY, CA,	CH, CN, CU, CZ,
DE, DK,	EE, ES, FI, GB,	GD, GE, GH, GM, HR,	HU, ID, IL, IN,
IS, JP,	KE, KG, KP, KR,	KZ, LC, LK, LR, LS,	LT, LU, LV, MD,
MG, MK,	MN, MW, MX, NO,	NZ, PL, PT, RO, RU,	SD, SE, SG, SI,
SK, SL,	TJ, TM, TR, TT,	UA, UG, US, UZ, VN,	YU, ZW, AM, AZ,
BY, KG,	KZ, MD, RU, TJ,	TM	
RW: GH, GM,	KE, LS, MW, SD,	SZ, UG, ZW, AT, BE,	CH, CY, DE, DK,
ES, FI,	FR, GB, GR, IE,	IT, LU, MC, NL, PT,	SE, BF, BJ, CF,
CG, CI,	CM, GA, GN, GW,	ML, MR, NE, SN, TD,	TG
AU 9917106	A1 19990705	AU 1999-17106	19981214
EP 1039926	A1 20001004	EP 1998-961904	19981214
R: AT, BE,	CH, DE, DK, ES,	FR, GB, GR, IT, LI,	LU, NL, SE, PT,
IE, FI			
PITY APPLN INFO	•	US 1997-68080	P 19971218

PRIORITY APPLN. INFO.:

US 1997-68080 P 19971218 WO 1998-US25725 W 19981214

The present invention describes a novel method of inducing immunity to cancer. This invention further discloses the use of Venezuelan Equine Encephalitis (VEE) virus vectors for expression of tumor-assocd. antigens, tumor-assocd. antigenic peptides and cytokines and methods for expressing these heterologous products in cultured cells, and in humans or animals.

REFERENCE COUNT:

9

REFERENCE(S):

(1) Brossart, P; J IMMUNOL 1997, V158, P3270

(3) Sidney, J; WO 9504542 A 1995 CAPLUS

(4) Sloan Kettering Inst Cancer; WO 9520974 A 1995 CAPLUS

- (5) Tuting, T; JOURNAL OF IMMUNOLOGY 1998, V160(3), P1139 CAPLUS
- (6) Tuting, T; JOURNAL OF MOLECULAR MEDICINE 1997, V75(7), P478 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 25 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1998:709094 CAPLUS

DOCUMENT NUMBER:

129:326928

TITLE:

Bifunctional proteins for cell-specific viral

vector targeting

CODEN: PIXXD2

INVENTOR(S):

Young, John; Snitkovsky, Sophie

PATENT ASSIGNEE(S):

President and Fellows of Harvard College, USA

SOURCE:

PCT Int. Appl., 74 pp.

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 9847916 A1 19981029 WO 1998-US7720 19980416

W: AU, CA, JP, NZ, US

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE

AU 9871277 A1 19981113 AU 1998-71277 19980416
PRIORITY APPLN. INFO.: US 1997-844359 19970418
WO 1998-US7720 19980416

AB The invention relates to a novel bifunctional mol. comprising a first binding moiety which binds to a surface mol. on a target cell and a second binding moiety which binds to a surface mol. on a viral vector. The bifunctional mol. targets the viral vector to the target cell with improved infectivity and selectivity. The mol. can be used, for example, in in vitro and in vivo gene delivery methods. Thus, a recombinant protein comprising epidermal growth factor fused to an Env protein-binding domain was used to target avian leukosis virus to EGF receptor-expressing cells.

L6 ANSWER 26 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1998:621324 CAPLUS

DOCUMENT NUMBER:

129:240848

TITLE:

Increasing the efficiency of uptake of

transforming DNA complexes with polycations

using peptides

Hawley-Nelson, Pamela; Lan, Jianqing; Shih, INVENTOR(S): Pojen; Jessee, Joel A.; Ciccarone, Valentina C.; Evans, Krista L.; Schifferli, Kevin P.; Gebeyehu, Guililat Life Technologies, Inc., USA PATENT ASSIGNEE(S): PCT Int. Appl., 105 pp. SOURCE: CODEN: PIXXD2 Patent DOCUMENT TYPE: LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. DATE PATENT NO. KIND DATE _____ _ _ _ _ WO 1998-US5232 19980316 WO 9840502 **A1** 19980917

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W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ,
             DE, DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP,
             KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK,
             MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL,
             TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ,
             MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES,
             FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG,
             CI, CM, GA, GN, ML, MR, NE, SN, TD, TG
                                                            19970314
                                           US 1997-818200
     US 6051429
                       Α
                            20000418
                            19980929
                                           AU 1998-65622
                                                            19980316
     AU 9865622
                      ` A1
                                           EP 1998-911737
                                                             19980316
     EP 1007699
                            20000614
                       A1
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC,
             PT, IE, FI
                                        US 1997-818200
                                                         A 19970314
PRIORITY APPLN. INFO.:
                                        US 1995-477354
                                                         B2 19950607
                                        US 1996-658130
                                                         A2 19960604
                                                         W 19980316
                                        WO 1998-US5232
```

AB A method of increasing the efficiency of transformation of eukaryotic cells using complexes of nucleic acids with polycations is decribed. The method uses peptide conjugates with nucleic acid-binding moieties, cationic lipids and dendrimers to complex the DNA. The peptides may be synthetic or derived from a cellular protein and may be further derivatized, e.g. by selective deprotection. The peptide may also be covalently linked to the transfection agent (lipid, cationic lipid or dendrimer). Inclusion of peptides or modified-peptides in transfection compns. or covalent attachment of peptides to transfection agents increases the efficiency of transfection. Methods for the prepn. of transfection compns. and methods of using these transfection compns. as intracellular delivery agents and extracellular targeting agents are also disclosed.

L6 ANSWER 27 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1998:381829 CAPLUS

DOCUMENT NUMBER: 129:134964

TITLE: Exogenous and endogenous IL-10 regulate

IFN-.alpha. production by peripheral blood

mononuclear cells in response to viral

stimulation

AUTHOR(S): Payvandi, Faribourz; Amrute, Sheela;

Fitzgerald-Bocarsly, Patricia

CORPORATE SOURCE: Dep. Pathol. Lab. Med., Graduate School Biomed.

Sci., Univ. Med. Dentistry/New Jersey, New Jersey Med. School, Newark, NJ, 07103, USA

SOURCE: J. Immunol. (1998), 160(12), 5861-5868

CODEN: JOIMA3; ISSN: 0022-1767

PUBLISHER: American Association of Immunologists

DOCUMENT TYPE: Journal LANGUAGE: English

IL-10 is an important regulator of the prodn. of proinflammatory AB cytokines. Its effect on IFN-.alpha. prodn., however, has not been reported. In this study, PBMC from healthy donors were stimulated with virus in the presence of IL-10. Human IL-10 (hIL-10) caused redns. in both the frequency of IFN-.alpha.-producing cells (IPC) and bulk IFN in response to herpes simplex virus type-1 (HSV-1), Sendai virus, Newcastle disease virus, and vesicular stomatitis The inhibitory effect occurred when IL-10 was added 2 or 4 h before, or 2 h poststimulation with HSV or Sendai virus, but not when added 4 h postinduction. Unlike IL-10, IL-4 did not affect the IFN-.alpha. response to HSV. However, when PBMC were induced with Sendai virus, IFN-.alpha. prodn. was also reduced by IL-4. treatment of PBMC resulted in strong redns. in the steady state levels of both HSV- and Sendai virus-induced IFN-.alpha.1, -.alpha.2, and -.beta. mRNA as detd. by RT-PCR. IFN-.alpha. prodn. of Sendai virus occurs predominantly by monocytes, whereas most enveloped viruses stimulate low frequency "natural IFN-producing cells (NIPC), " which are thought to be dendritic cells. Peripheral blood dendritic cells were found to express the IL-10 receptor, suggesting that IL-10 may directly act on the dendritic IPC. Addn. of monoclonal anti-IL-10 to PBMC resulted in a significant increase in both the frequency of IPC and the amt. of secreted IFN-.alpha. in response to HSV but not Sendai virus. We conclude that human IL-10 can serve as both an endogenous and exogenous regulator of IFN-.alpha. prodn.

L6 ANSWER 28 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1996:643186 CAPLUS

DOCUMENT NUMBER: 125:324125

TITLE: The .beta.-amyloid domain is essential for axonal sorting of amyloid precursor protein

AUTHOR(S): Tienari, Pentti J.; De Strooper, Bart; Ikonen,

Elina; Simons, Mikael; Weidemann, Andreas;

Czech, Christian; Hartmann, Tobias; Ida, Nobuo;

Multhaup, Gerd; et al.

CORPORATE SOURCE: Cell Biol. Programme, European Molecular Biology

Lab., Heidelberg, D-69012, Germany

SOURCE: EMBO J. (1996), 15(19), 5218-5229

CODEN: EMJODG; ISSN: 0261-4189

DOCUMENT TYPE: Journal LANGUAGE: English

The authors have analyzed the axonal sorting signals of amyloid AB precursor protein (APP). Wild-type and mutant versions of human APP were expressed in hippocampal neurons using the Semliki forest virus system. The authors show that wild-type APP and mutations implicated in Alzheimer's disease and another brain .beta.-amyloidosis are sorted to the axon. By anal. of deletion mutants the authors found that the membrane-inserted APP ectodomain but not the cytoplasmic tail is required for axonal sorting. Systematic deletions of the APP ectodomain identified two regions required for axonal delivery: one encoded by exons 11-15 in the carbohydrate domain, the other encoded by exons 16-17 in the juxtamembranous .beta.-amyloid domain. Treatment of the cells with the N-glycosylation inhibitor tunicamycin induced missorting of wild-type APP, supporting the importance of glycosylation in axonal sorting of APP. The data revealed a hierarchy of sorting signals on APP: the .beta.-amyloid-dependent membrane proximal signal was the major contributor to axonal sorting, while N-glycosylation had a weaker effect. Furthermore, recessive somatodendritic signals, most likely in the cytoplasmic tail, directed the protein to the dendrites when the ectodomain was deleted. Anal. of delivered protein, hemagglutinin, demonstrated that only hemagglutinin formed CHAPS-insol. complexes, suggesting distinct mechanisms of axonal sorting for these two proteins. This study is the first delineation of sorting requirements of an axonally targeted protein in polarized neurons and indicates that the .beta.-amyloid domain plays a major role in axonal delivery of APP.

L6 ANSWER 29 OF 37 CAPLUS COPYRIGHT 2001 ACS

DOCUMENT NUMBER: 125:164939

ACCESSION NUMBER:

TITLE: Expression and analysis of presentlin 1 in a

human neuronal system: Localization in cell

bodies and dendrites

1996:524456 CAPLUS

AUTHOR(S): Cook, David G.; Sung, Jane C.; Golde, Todd E.;

Felsenstein, Kevin M.; Wojczyk, Boguslaw S.; Tanzi, Rudolph E.; Trojanowski, John Q.; Lee,

Virginia M.-Y.; Doms, Robert W.

CORPORATE SOURCE: Dep. Pathology, Univ. Pennsylvania,

Philadelphia, PA, 19104, USA

SOURCE: Proc. Natl. Acad. Sci. U. S. A. (1996), 93(17),

9223-9228

CODEN: PNASA6; ISSN: 0027-8424

DOCUMENT TYPE: Journal LANGUAGE: English

Mutations in the recently identified presentlin 1 gene on chromosome AB 14 cause early onset familial Alzheimer disease (FAD). Here, the author's describe the expression and anal. of the protein coded by presenilin 1 (PS1) in NT2N neurons, a human neuronal model system. PS1 was expressed using recombinant Semliki Forest virions and detected by introduced antigenic tags or antisera to PS1-derived peptides. Immunopptn. revealed 2 major PS1 bands of approx. 43 and 50 kDa, neither of which were N-glycosylated or O-glycosylated. Immunoreactive PS1 was detected in cell bodies and dendrites of NT2N neurons but not in axons or on the cell surface. PS1 was also detected in BHK cells, where it was also intracellular and colocalized with calnexin, a marker for the rough endoplasmic reticulum. A mutant form of PS1 linked to FAD did not differ from the wild-type protein at the light microscopic level. The model system described here will enable studies of the function of PS1 in human neurons and the role of mutant PS1 in FAD.

L6 ANSWER 30 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1996:358345 CAPLUS

DOCUMENT NUMBER: 125:31828

TITLE: Phenotypic changes in Langerhans' cells after

infection with arboviruses: a role in the immune

response to epidermally acquired viral

infection?

AUTHOR(S): Johnston, Linda J.; Halliday, Gary M.; King,

Nicholas J. C.

CORPORATE SOURCE: Dep. Pathol. Med., Univ. Sydney, Sydney, 2006,

Australia

SOURCE: J. Virol. (1996), 70(7), 4761-4766

CODEN: JOVIAM; ISSN: 0022-538X

DOCUMENT TYPE: Journal LANGUAGE: English

The role of Langerhans cells (LC) in the initiation of an immune response to a viral infection remains unclear. In vivo epidermal infection with the arboviruses West Nile virus and Semliki Forest virus significantly increased the expression of major histocompatibility complex class II antigens, CD54, and CD80 on LC. Thus, during an epidermally acquired viral infection, local LC appear to mature to a phenotype approximating that of lymphoid dendritic cells. This change may be important in the activation of naive T cells and the subsequent clearance of viral infection.

L6 ANSWER 31 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1995:797180 CAPLUS

DOCUMENT NUMBER:

123:196271

TITLE:

Intracellular routing of wild-type and mutated polymeric immunoglobulin receptor in hippocampal

neurons in culture

AUTHOR(S):

de Hoop, Meltsje; von Poser, Christine; Lange, Claudia; Ikonen, Elina; Hunziker, Walter; Dotti,

Carlos G.

CORPORATE SOURCE:

Cell Biology Program, European Molecular Biology

Laboratory, Heidelberg, 69012, Germany

SOURCE:

J. Cell Biol. (1995), 130(6), 1447-59

CODEN: JCLBA3; ISSN: 0021-9525

DOCUMENT TYPE:

Journal English

LANGUAGE:

AB

Certain epithelial cells synthesize the polymeric Ig receptor (pIgR) to transport IgA and IgM into external secretions. In polarized epithelia, newly synthesized receptor is first delivered to the basolateral plasma membrane and is then, after binding the Ig,

transcytosed to the apical plasma membrane, where the receptor-ligand complex is released by proteolytic cleavage. In a previous work (E. Ikonen, et al., 1993), the authors implied the

existence of a dendro-axonal transcytotic pathway for the rabbit pIgR expressed in hippocampal neurons via the **Semliki Forest** virus (SFV) expression system. By labeling

surface-exposed pIgR in live neuronal cells, the authors now show

(a) internalization of the receptor from the dendritic plasma membrane to the dendritic early endosomes, (b) redistribution of the receptor from the dendritic to the

axonal domain, (c) inhibition of this movement by brefeldin A, (BFA) and (d) stimulation by the activation of protein kinase C (PKC) via phorbol myristate acetate (PMA). In addn., the authors show that a mutant form of the receptor lacking the epithelial basolateral sorting signal is directly delivered to the axonal domain of hippocampal neurons. Although this mutant is internalized into early endosomes, no transcytosis to the dendrites could be

early endosomes, no transcytosis to the **dendrites** could be obsd. In epithelial Madin-Darby Canine kidney (MDCK) cells, the mutant receptor could also be internalized into basolaterally derived early endosomes. These results suggest the existence of a dendro-axonal transcytotic pathway in neuronal cells which shares

epithelial cells.

ANSWER 32 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1995:622960 CAPLUS

similarities with the basolateral to apical transcytosis in

DOCUMENT NUMBER:

123:53.039

TITLE:

L6

Nocodazole-dependent transport, and brefeldin

A-sensitive processing and sorting, of newly synthesized membrane proteins in cultured

neurons

Cid-Arregui, Angel; Parton, Robert G.; Simons, AUTHOR (S):

Kai; Dotti, Carlos G.

Cell Biol. Prog., European Mol. Biol. Lab., CORPORATE SOURCE:

Heidelberg, D-69012, Germany

J. Neurosci. (1995), 15(6), 4259-69 SOURCE:

CODEN: JNRSDS; ISSN: 0270-6474

Journal DOCUMENT TYPE:

English LANGUAGE:

> The envelope glycoproteins of Semliki Forest virus (SFV), Vesicular Stomatitis virus (VSV), and Influenza Fowl Plaque virus (FPV) are vectorially targeted in neurons to the plasma membrane of dendrites (SFV and VSV) and axons (FPV). gain insight into the mechanisms responsible for such polarized delivery we have examd. the effects on neurons of nocodazole and brefeldin A (BFA), which are known to cause microtubule depolymn. and disassembly of the Golgi app., resp. Nocodazole treatment blocked transport of all viral glycoproteins to both axons and dendrites. BFA treatment induced disruption of the Golgi complex, including the trans-Golgi network (TGN), and tubulation of endosomes. However, the delivery of the SFV and FPV glycoproteins to the cell surface was not affected significantly by BFA, although processing and sorting were altered, as revealed by surface biotinylation and immunofluorescence microscopy of fixed nonpermeabilized cells. These results demonstrate the involvement of microtubules in axonal and dendritic transport of integral membrane glycoproteins, and the existence of a BFA-sensitive component in the sorting but not in the transport machinery.

L6 ANSWER 33 OF 37 CAPLUS COPYRIGHT 2001 ACS

1995:548456 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 122:311663

Intracellular routing of human amyloid protein TITLE:

precursor: axonal delivery followed by transport

to the dendrites

Simons, M.; Ikonen, E.; Tienari, P. J.; AUTHOR(S):

Cid-Arregui, A.; Moenning, U.; Beyreuther, K.;

Dotti, C. G.

CORPORATE SOURCE: Cell Biology Program, Univ. of heidelberg,

Heidelberg, Germany

J. Neurosci. Res. (1995), 41(1), 121-8 SOURCE:

CODEN: JNREDK; ISSN: 0360-4012

DOCUMENT TYPE: Journal

English LANGUAGE:

feature of Alzheimer's disease is the A characteristic neuropathol.

cerebral deposition of amyloid plaques. These deposits contain .beta.A4 amyloid peptide, a cleavage product of the transmembrane protein amyloid protein precursor (APP). Despite numerous studies on the processing of the different APP isoforms in non-neuronal cells, little is known about its sorting and transport in neurons of the central nervous system (CNS). To analyze this question the authors expressed in cultured rat hippocampal neurons the human APP 695, tagged at its N-terminus with he myc epitope, using the Semliki forest virus (SFV) expression system. APP was first delivered from the cell body to the axon and later appeared also in the dendrites. Inhibition of protein synthesis at the time of axonal expression did not block the late appearance of the protein in the dendrites. An antibody directed against the myc tag, bound to the cell surface at 4.degree.C at the time of axonal APP expression, could be chased to the dendritic domain after subsequent incubation at 37.degree.C. These results suggest that the newly synthesized APP, after initial axonal delivery, may be transported to the dendrites by a transcytotic mechanism. The routing of APP in polarized neurons is different from that of polarized epithelial cells, in which the protein is delivered basolaterally, arguing for neuronal specific sorting and processing mechanisms.

ANSWER 34 OF 37 CAPLUS COPYRIGHT 2001 ACS L6

1994:6446 CAPLUS ACCESSION NUMBER:

120:6446 DOCUMENT NUMBER:

Transcytosis of the polymeric immunoglobulin TITLE:

receptor in cultured hippocampal neurons

Ikonen, Elina; Parton, Robert G.; Hunziker, AUTHOR(S):

Walter; Simons, Kai; Dotti, Carlos G.

Cell Biol. Program, Eur. Mol. Biol. Lab., CORPORATE SOURCE:

Heidelberg, D-69012, Germany

Curr. Biol. (1993), 3(10), 635-44 SOURCE:

CODEN: CUBLE2; ISSN: 0960-9822

Journal DOCUMENT TYPE: English LANGUAGE:

AB The authors report expression of the polymeric Ig receptor in cultured hippocampal neurons, using a Semliki Forest Virus expression system, and show by immunofluorescence microscopy that the newly synthesized receptor is targeted from the Golgi complex predominantly to the dendrites only; .apprx.20% of the infected neurons display axonal immunofluorescence. Addn. of ligand leads to significant redistribution of the receptor to the axons, shown by an .apprx.3 fold increase in axonal immunoreactivity with the anti-receptor antibodies. Thus, a transcytotic route, analogous to that in epithelia, exists in neurons, where it transports proteins from the

somatodendritic to the axonal domain. Cultured neurons expressing

the polymeric Ig receptor offer an exptl. system that should be useful for further characterization of this novel neuronal pathway at the mol. and functional level.

L6 ANSWER 35 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1993:600304 CAPLUS

DOCUMENT NUMBER: 119:200304

TITLE: Protein transport to the dendritic

plasma membrane of cultured neurons is regulated

by rab8p

AUTHOR(S): Huber, Lukas A.; de Hoop, Meltsje J.; Dupree,

Paul; Zerial, Marino; Simons, Kai; Dotti, Carlos

CORPORATE SOURCE: Cell Biol. Programme, Eur. Mol. Biol. Lab.,

Heidelberg, D-69012, Germany

SOURCE: J. Cell Biol. (1993), 123(1), 47-55

CODEN: JCLBA3; ISSN: 0021-9525

DOCUMENT TYPE: Journal LANGUAGE: English

In the companion paper (Huber, L. A.; et al., 1993) the authors AB reported that the small GTPase rab8p is involved in transport from the TGN to the basolateral plasma membrane in epithelia. In the present work the authors investigated the localization and function of rab8p in polarized hippocampal neurons. By immunofluorescence microscopy the authors found that rab8p localized preferentially in the somatodendritic domain, and was excluded from the axon. Double-labeling immunofluorescence showed that some of the rab8p colocalized in the dendrites with the Semliki Forest Virus qlycoprotein E2 (SFV-E2). An antisense oligonucleotide approach was used to investigate the role of rab8p in dendritic transport of newly synthesized viral glycoproteins. Antisense oligonucleotides corresponding to the initiation region of the rab8 coding sequence were added to the cultured neurons for 4 days. This treatment resulted in a significant decrease in cellular levels of rab8p and transport of SFV-E2 from the cell body to the dendrites was significantly reduced. However, no effect was obsd. on axonal transport of influenza HA. Thus, rab8p is involved in transport of proteins to the dendritic surface in neurons.

L6 ANSWER 36 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1993:512568 CAPLUS

DOCUMENT NUMBER: 119:112568

TITLE: Expression of heterologous proteins in cultured

rat hippocampal neurons using the

Semliki Forest virus vector

AUTHOR(S): Olkkonen, V. M.; Liljestroem, P.; Garoff, H.;

Simons, K.; Dotti, C. G.

CORPORATE SOURCE: Cell Biol. Programme, Eur. Mol. Biol. Lab.,

Heidelberg, 6900, Germany

SOURCE: J. Neurosci. Res. (1993), 35(4), 445-51

CODEN: JNREDK; ISSN: 0360-4012

DOCUMENT TYPE: Journal LANGUAGE: English

AB The Semliki Forest virus expression vector

(Liljestroem, P.; Garoff, H., 1991) was tested in cultured rat hippocampal neurons using two MDCK cell membrane-assocd. proteins as reporters: rab8, a small GTPase involved in post-Golgi vesicle transport, and VIP21, an integral membrane protein of caveolae, trans-Golgi network, and post-Golgi vesicles. Expression of the c-myc epitope-tagged proteins was visualized by immunofluorescence microscopy. The proteins were first detected in neurons after 3-4 h infection by the recombinant viruses. The infection efficiency on neurons was high: after 6 h infection at a multiplicity of one, 50-60% of the cells expressed the reporter proteins. The neurons tolerated the infection well .ltoreq.8 h. Their polarized organization was not disturbed, as judged from morphol. and from distribution of the dendritic MAP2 and axonal synaptophysin marker proteins. The Semliki Forest virus vector thus seems suitable for short-term expression of proteins in cultured neurons.

L6 ANSWER 37 OF 37 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1993:405888 CAPLUS

DOCUMENT NUMBER: 119:5888

TITLE: Polarized distribution of the viral

glycoproteins of vesicular stomatitis, fowl

plaque and Semliki Forest

viruses in hippocampal neurons in culture: a

light and electron microscopy study

AUTHOR(S): Dotti, Carlos G.; Kartenbeck, Juergen; Simons,

Kai

CORPORATE SOURCE: Cell Biology Program, European Molecular Biology

Laboratory, Heidelberg, Germany Brain Res. (1993), 610(1), 141-7

CODEN: BRREAP; ISSN: 0006-8993

DOCUMENT TYPE: Journal LANGUAGE: English

SOURCE:

AB It has been shown previously using immunofluorescence microscopy that upon infection of polarized hippocampal cells in culture with vesicular stomatitis virus (VSV) and fowl plague virus (FPV) the VSV glycoprotein is delivered to the plasma membrane of the dendrites and of the cell body, whereas the FPV hemagglutinin is transported to the axonal surface. In this work electron microscopy of infected rat hippocampal neurons showed that VSV progeny budded from the plasma membrane of the dendrites and the cell body. The location of the budding virions corresponded

to the distribution of the VSV glycoprotein which was detected over the somatodendritic plasma membrane by immunoelectron microscopy. In contrast, no FPV formation was seen in the infected neurons, although the FPV hemagglutinin was localized to the axonal surface by immunoelectron microscopy. In Semliki Forest virus (SFV)-infected hippocampal cells, the viral glycoproteins were exclusively present in the dendrites and cell body but not in axons.

(FILE MEDILINE, BIOSIS, EMBASE, WPIDS, CONFSCI, SCISEARCH, JICST-EPLUS, JAPIO' ENTERED AT 11:40:47 ON 19 SEP 2001)

L7 136 S L6

 $rac{1}{8}$

13 S L7 AND ("E2" OR RESIDUE(5A)160)

L9 45 S L7 AND RECOMBINAN?

57 S L8 OR L9

33 Dup Rum L10 (24 Duplicates Removed)

L11 ANSWER 1 OF 33 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER: 2001-316326 [33] WPIDS

CROSS REFERENCE: 2001-308587 [31]

DOC. NO. CPI: C2001-097446

TITLE: New isolated and purified gp100 useful for the

prophylactic treatment of cancer.

DERWENT CLASS: B04 D16

INVENTOR(S): BARBER, B; BERINSTEIN, N; MOINGEON, P; TARTAGLIA,

J; TINE, J A

PATENT ASSIGNEE(S): (AVET) AVENTIS PASTEUR LTD

COUNTRY COUNT: 94

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2001030847 A1 20010503 (200133) * EN 89

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE

DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG

KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ

PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2001010137 A 20010508 (200149)

APPLICATION DETAILS:

PATENT NO K	CIND	API	PLICATION	DATE
WO 2001030847	' A1	WO	2000-CA1254	20001020
AU 2001010137	' A	AU	2001-10137	20001020

FILING DETAILS:

PRIORITY APPLN. INFO: US 2000-223325 20000807; US 1999-160879

19991022

AN 2001-316326 [33] WPIDS

CR 2001-308587 [31]

AB WO 200130847 A UPAB: 20010831

NOVELTY - An isolated and purified modified gp100 molecule (N1) capable of modulating an immune response in an animal is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are included for:

- (1) a cell comprising (N1) where the cell expresses a polypeptide encoded by the nucleic acid;
- (2) a recombinant virus comprising a virus into which(N1) has been inserted which encodes for a polypeptide, where the virus causes the expression of the polypeptide in an infected cell;
- (3) a recombinant virus into which (N1) has been inserted which encodes for a polypeptide where cells infected with the virus are capable of eliciting an immune response directly against a member selected from:
 - (a) the polypeptide;
 - (b) a fragment of the polypeptide;
 - (c) a cell expressing the polypeptide or a fragment of it; or
 - (d) cells binding the protein or fragment of it;
 - (4) an isolated protein encoded by (N1);
- (5) an isolated protein having the activity of a modified gp100 protein;
- (6) a protein having a sequence of 661 amino acids described in the specification;
- (7) modulating an animal's immune system comprising administering an effective amount of a gp100 or gp100 protein which has been modified;
- (8) modulating an animal's immune system comprising administering to an animal in need of it, an effective amount of a vector, into which gp100 which has been modified is inserted;
- (9) prophylactic treatment of cancer comprising administering to an animal an effective amount of a modified gp100 or immunogenic fragment of it, or a nucleic acid sequence encoding a modified gp100 or immunogenic fragment of it;
- (10) a melanoma vaccine comprising a nucleic acid sequence encoding a modified gp100;
- (11) a modified gp100 protein sequence which is modified to enhance its binding to MHC molecules;
 - (12) a vaccine comprising a modified gp100 nucleic acid

sequence or its corresponding protein or protein fragment capable of eliciting the production of antibodies in a animal to corresponding antigens;

- (13) a vaccine comprising a modified gp100 nucleic acid sequence or its corresponding protein or protein fragment capable of eliciting a cellular immune response; and
- (14) an immunogenic composition containing a vaccine vector encoding for a modified gp100 molecule.

ACTIVITY - Cytostatic.

MECHANISM OF ACTION - Vaccine; Gene therapy.

USE - Nucleic acids and proteins of the invention are useful as vaccines for prophylactic treatment of cancer (claimed).

Dwg.0/12

L11 ANSWER 2 OF 33 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER:

2001-235115 [24] WPIDS

DOC. NO. CPI:

C2001-070478

TITLE:

Introducing nucleic acid into cell, useful e.g. in

gene therapy of tumors, using alphavirus

vector and vector-specific antibody to enhance

infection.

DERWENT CLASS:

B04 C06 D16

INVENTOR (S):

JOHNSTON, R E; MACDONALD, G H (UYNC-N) UNIV NORTH CAROLINA

COUNTRY COUNT:

94

PATENT INFORMATION:

PATENT ASSIGNEE(S):

PATENT N	IO	KIND	DATE	WEEK	LA	PG
			. 	 -		-

WO 2001016343 A1 20010308 (200124) * EN 66

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW MZ NL OA PT SD SE SL SZ TZ UG ZW

W: AE AG AL AM AT AU AZ BA BB BG BR BY BZ CA CH CN CR CU CZ DE DK DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX MZ NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA ZW

AU 2000074718 A 20010326 (200137)

APPLICATION DETAILS:

	KIND	APPLICATION	DATE
WO 200101634		WO 2000-US23845	
AU 200007471	18 A	AU 2000-74718	20000830

FILING DETAILS:

PATENT NO KIND

PATENT NO

AU 2000074718 A Based on

WO 200116343

PRIORITY APPLN. INFO: US 2000-177435 20000121; US 1999-151718 19990831

AN 2001-235115 [24] WPIDS

AB WO 200116343 A UPAB: 20010502

NOVELTY - Introducing and expressing a nucleic acid (I) in a cell by contacting the cell with an **alphavirus** vector (A) containing heterologous (I), and an antibody (Ab) that binds specifically to (A), is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) delivering (I) to a subject by administering:
- (a) (A) and Ab;
 - (b) (A) only if the subject already contains Ab; or
 - (c) a cell previously contacted with (A) and Ab;
 - (2) generating an immune response, by administering:
 - (a) an (A) in which (I) encodes an immunogen and Ab;
 - (b) (A) only if the subject already contains Ab; or
 - (c) a cell previously contacted with (A) and Ab; and
- (3) pharmaceutical formulation comprising a complex of (A) and Ab in a carrier.

ACTIVITY - Cytostatic; antiviral; antibacterial; antiprotozoal; antiparasitic.

No biological data is given.

MECHANISM OF ACTION - Induction of specific immune responses, both humoral and cellular; gene therapy; low concentrations of Ab form, with (A), complexes that interact with Fc or complement receptors on cells, preferably APCs, resulting in increased infectivity.

USE - The method is used to express (I) in humans or other animals (mammals and birds). (I) encodes an antigen from a tumor or infectious agent, particularly a latent or chronic agent, including bacterium, virus, protozoan or parasite. (I) may encode a therapeutic protein, for treating a wide range of genetic or acquired disorders, a protein that confers resistance to anticancer agents or an immunostimulant. Alternatively, (I) encodes a therapeutic RNA. (All claimed). (I) can be used to treat infectious diseases, or tumors. The method can be used for transient or stable expression of (I) in cultured cells, or for preparation of transgenic animals. Antibodies raised against the new immunogenic compositions can be used for passive immunization or for diagnosis or histological applications.

ADVANTAGE - Ab increases the infectivity of (A) for particular cell types and may target (A) to antigen-presenting cells, resulting in an increased immune response, i.e. even weak antigens that fail

when used in conventional compositions may be used successfully. When formulated with Ab, (A) induces a stronger immune response than the corresponding infectious organism itself. With alphaviruses, antibody-dependent enhancement is not associated with significant pathology.

Dwg.0/8

L11 ANSWER 3 OF 33 MEDLINE

ACCESSION NUMBER: 2001231630 MEDLINE

DOCUMENT NUMBER: 21221117 PubMed ID: 11296257

TITLE: Activation-dependent changes in receptor distribution

and dendritic morphology in hippocampal

neurons expressing P2X2-green fluorescent protein

receptors.

AUTHOR: Khakh B S; Smith W B; Chiu C S; Ju D; Davidson N;

Lester H A

CORPORATE SOURCE: Division of Biology, 156-29, California Institute of

Technology, Pasadena, CA 91125, USA...

bsk@mrc-lmb.com.ac.uk

CONTRACT NUMBER: MH49176 (NIMH)

NS-11756 (NINDS)

SOURCE: PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF

THE UNITED STATES OF AMERICA, (2001 Apr 24) 98 (9)

5288-93.

Journal code: PV3; 7505876. ISSN: 0027-8424.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200105

ENTRY DATE: Entered STN: 20010529

Last Updated on STN: 20010529 Entered Medline: 20010521

ATP-gated P2X(2) receptors are widely expressed in neurons, but the cellular effects of receptor activation are unclear. We engineered functional green fluorescent protein (GFP)-tagged P2X(2) receptors and expressed them in embryonic hippocampal neurons, and report an approach to determining functional and total receptor pool sizes in living cells. ATP application to dendrites caused receptor redistribution and the formation of varicose hot spots of higher P2X(2)-GFP receptor density. Redistribution in dendrites was accompanied by an activation-dependent enhancement of the ATP-evoked current. Substate-specific mutant T18A P2X(2)-GFP receptors showed no redistribution or activation-dependent enhancement of the ATP-evoked current. Thus fluorescent P2X(2)-GFP receptors function normally, can be quantified, and reveal the dynamics of P2X(2) receptor distribution on the seconds time scale.

L11 ANSWER 4 OF 33 SCISEARCH COPYRIGHT 2001 ISI (R)

2001:391128 SCISEARCH ACCESSION NUMBER:

THE GENUINE ARTICLE: 430EG

Sindbis viral-mediated expression of TITLE:

> Ca2+-permeable AMPA receptors at hippocampal CA1 synapses and induction of NMDA receptor-independent

long-term potentiation

Okada T; Yamada N; Kakegawa W; Tsuzuki K; Kawamura **AUTHOR:**

M; Nawa H; Iino M; Ozawa S (Reprint)

Gunma Univ, Sch Med, Dept Physiol, 3-39-22 Showa CORPORATE SOURCE:

> Machi, Gunma 3718511, Japan (Reprint); Gunma Univ, Sch Med, Dept Physiol, Gunma 3718511, Japan; Japan

Sci & Technol Corp, CREST, Kawaguchi, Saitama

3320012, Japan; Niigata Univ, Brain Res Inst, Dept

Mol Neurobiol, Niigata 9518585, Japan

COUNTRY OF AUTHOR:

Japan

SOURCE:

EUROPEAN JOURNAL OF NEUROSCIENCE, (APR 2001) Vol.

13, No. 8, pp. 1635-1643.

Publisher: BLACKWELL SCIENCE LTD, P O BOX 88, OSNEY

MEAD, OXFORD OX2 ONE, OXON, ENGLAND.

ISSN: 0953-816X. Article; Journal

DOCUMENT TYPE:

LANGUAGE:

English

REFERENCE COUNT:

34

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Gene manipulation in order to artificially express a particular AB gene in neurons in the central nervous system is a powerful tool for the analysis of brain function. Sindbis viral vectors have been developed to express high levels of foreign genes in postmitotic brain neurons with little transfection of glial cells. In this study, we expressed the gene encoding the unedited GluR2 (GluR-B) subunit of the AMPA-type glutamate receptor that forms inwardly rectifying and Ca2+-permeable channels, in rat CA1 hippocampal neurons in slice cultures using Sindbis viral vectors. The pyramidal cell layer of the CA1 region was injected with recombinant Sindbis viruses encoding both enhanced green fluorescent protein (GFP) and unedited GluR2. The GFP fluorescence from CA1 neurons could be detected as early as 6 h and reached a maximal level about 48 h postinfection. The inwardly rectifying and Ca2+-permeable AMPA receptors were expressed in most CA1 pyramidal cells expressing GFP. These AMPA receptors expressed by gene transfer were involved in fast excitatory neurotransmission elicited by electrical stimulation of the Schaffer collaterals in the stratum radiatum. Tetanic stimulation of Schaffer collaterals induced NMDA receptor-independent, long-term potentiation due to Ca2+ influx through the newly expressed AMPA receptors in the area densely stained with GFP. Thus, the combined use of Sindbis viral vectors with the GFP reporter allowed physiological

examination of the roles of a specific gene product in synaptic function in well-characterized brain neurons.

L11 ANSWER 5 OF 33 MEDLINE DUPLICATE 1

ACCESSION NUMBER: 2001450312 IN-PROCESS

DOCUMENT NUMBER: 21387414 PubMed ID: 11495971

TITLE: Abundant gfp expression and ltp in hippocampal acute

slices by in vivo injection of sindbis

virus.

AUTHOR: D'Apuzzo M; Mandolesi G; Reis G; Schuman E M

CORPORATE SOURCE: California Institute of Technology, Pasadena,

California 91125.

SOURCE: JOURNAL OF NEUROPHYSIOLOGY, (2001 Aug) 86 (2)

1037-42.

Journal code: JC7; 0375404. ISSN: 0022-3077.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: IN-PROCESS; NONINDEXED; Priority Journals

ENTRY DATE: Entered STN: 20010813

Last Updated on STN: 20010813

Virus-mediated gene transfer into neurons is a powerful tool for the AB analysis of neuronal structure and function. Recombinant sindbis virus has been previously used to study protein function in hippocampal neuron cultures as well as in hippocampal organotypic slice cultures. Nevertheless, some concern still exists about the physiological relevance of these cultured preparations. Acute hippocampal slices are a widely used preparation for the study of synaptic transmission, but currently recombinant gene delivery is usually achieved only through time-consuming transgenic techniques. In this study, we show that a subregion of the CA1 area in acute hippocampal slices can be specifically altered to express a gene of interest. A sindbis virus vector carrying an enhanced green fluorescent protein (EGFP) reporter was injected in vivo into the hippocampus of adult rats. After 18 h, rats were killed, and acute hippocampal slices, infected in the CA1 field, were analyzed morphologically and electrophysiologically. Infected slices showed healthy and stable electrophysiological responses as well as long-term potentiation. In addition, infected pyramidal cells were readily recognized in living slices by two-photon imaging. Specifically, the introduction of an EGFP-Actin fusion protein greatly enhanced the detection of fine processes and dendritic spines. We propose this technique as an efficient tool for studying gene function in adult hippocampal neurons.

L11 ANSWER 6 OF 33 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 2001290084 EMBASE

TITLE: Gene delivery of vaccines for infectious disease.

Clark K.R.; Johnson P.R. **AUTHOR:**

P.R. Johnson, Children's Research Institute, CORPORATE SOURCE:

Department of Pediatrics, The Ohio State University,

Columbus, OH 43205, United States. johnsonp@pediatrics.ohio-state.edu

Current Opinion in Molecular Therapeutics, (2001) 3/4 SOURCE:

> (375-384). Refs: 65

ISSN: 1464-8431 CODEN: CUOTFO

COUNTRY: DOCUMENT TYPE:

United Kingdom Journal; Article 004 Microbiology FILE SEGMENT: 022 Human Genetics

> 026 Immunology, Serology and Transplantation

029 Clinical Biochemistry 037 Drug Literature Index

039 Pharmacy

LANGUAGE: English English SUMMARY LANGUAGE:

Genetic immunization is the process of delivering and expressing a AB gene (or therapeutic nucleic acid) encoding a pathogen-derived antigen into target host cells to elicit a protective humoral or cell-mediated immune response. Gene delivery methods to achieve this goal have expanded rapidly, and currently employ a variety of oligonucleotides, synthetic polypeptides, recombinant vectors and even edible plants, all of which have been shown to be capable of inducing protective immunity in experimental animal models. This review highlights recent progress in several gene delivery systems (both non-viral and viral methods) using novel in vivo approaches to engender effective host immune responses against the introduced antigen.

L11 ANSWER 7 OF 33 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.

ACCESSION NUMBER: 2001081556 EMBASE

Web alert: Molecular vaccines for disease prevention TITLE:

and therapy.

AUTHOR: Jones D.

D. Jones, Current Drugs Ltd, Middlesex House, 34-42 CORPORATE SOURCE:

Cleveland St., London W1T 4LB, United Kingdom.

daniel.jones@current-drugs.com

Current Opinion in Molecular Therapeutics, (2001) 3/1 SOURCE:

(11-12).

ISSN: 1464-8431 CODEN: CUOTFO

United Kingdom COUNTRY: Journal; Article DOCUMENT TYPE:

Microbiology FILE SEGMENT: 004

> 016 Cancer

026 Immunology, Serology and Transplantation

> Shears 308-4994 Searcher :

037 Drug Literature Index

LANGUAGE: English

L11 ANSWER 8 OF 33 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER:

2000-619231 [59] WPIDS

DOC. NO. CPI:

C2000-185549

TITLE:

New alphavirus that infects human

dendritic cells for use in generating an immune response to pathogenic agents such as

bacteria, viruses, fungi, parasites and cancer and

for biological assays.

DERWENT CLASS:

B04 D16

INVENTOR(S):

BARNETT, S; DRIVER, D A; DUBENSKY, T W; FROLOV, I;

GARDNER, J P; OTTEN, G; POLO, J M

PATENT ASSIGNEE(S):

(CHIR) CHIRON CORP

COUNTRY COUNT:

92

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2000061772 A2 20001019 (200059)* EN 83

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ TZ UG ZW

 W: AE AG AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK

 DM DZ EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP

 KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX NO NZ PL PT

 RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG US UZ VN YU ZA

zw

AU 2000043660 A 20001114 (200108)

APPLICATION DETAILS:

PATENT NO KIND	APPLICATION	DATE
WO 2000061772 A2	WO 2000-US10722	20000414
AU 2000043660 A	AU 2000-43660	20000414

FILING DETAILS:

PATENT NO KIND PATENT NO

AU 2000043660 A Based on WO 200061772

PRIORITY APPLN. INFO: US 2000-191363 20000322; US 1999-129498 19990414; US 1999-148086 19990809

AN 2000-619231 [59] WPIDS

AB WO 200061772 A UPAB: 20001117

NOVELTY - An isolated alphavirus (AV) which infects human

dendritic cells and is not of American Type Culture Collection (ATCC) number VR-2526, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) an isolated AV which infects non-human dendritic cells and is not a Venezuelan equine encephalitis virus or ATCC VR-2526;
- (2) an isolated nucleic acid comprising a nucleic acid which encodes AV;
- (3) an isolated nucleic acid comprising a nucleic acid that encodes an AV, having a sequence of 11703 nucleotides, given in the specification;
- (4) an AV structural protein cassette, comprising a promoter operably linked to a nucleic acid sequence encoding AV structural proteins (sP) from the new AV;
 - (5) an AV packaging cell, comprising a host cell and (4);
- (6) an AV producer cell, comprising (5) and an alphavirus RNA vector replicon, an alphavirus vector construct, or a eukaryotic layered vector initiation system;
- (7) a **recombinant** AV particle, comprising a particle produced from a cell line of (6);
- (8) a **recombinant** AV particle, comprising a particle produced from a cell line of (5);
- (9) a recombinant AV particle which infects human dendritic cells and that is not derived from ATCC VR -2526;
- (10) a recombinant AV particle which infects non-human dendritic cells and that is not derived from a Venezuelan equine encephalitis virus or ATCC VR-2526;
- (11) introducing a heterologous nucleotide sequence into cells, comprising infecting the cells with (7), (8), (9) or (10);
 - (12) an AV vector construct comprising:
- (a) a 5' promoter that initiates synthesis of viral RNA in vitro from cDNA;
 - (b) a 5' sequence that initiates transcription of AV RNA;
- (c) a nucleic acid molecule that operably encodes all 4 AV
 nsP's;
 - (d) an AV RNA polymerase recognition sequence; and
- (e) a 3' polyadenylate tract, where the nucleic acid sequence that operably encodes all 4 AV nonstructural proteins (nsP) contains a mutation in at least one nsP that is a mutation in nsP1 residues 346, 441, 473, nsP2 residues 438, 622, 634, 715, nsP3 residues, 417, 456, 505, and nsP4 residue 266, as compared to wild-type;
- (13) a eukaryotic layered vector initiation system, comprising a 5' promoter capable of initiating in vivo the 5' synthesis of AV RNA from cDNA, a sequence which initiates transcription of AV RNA following the 5' promoter, a nucleic acid which operably encodes all

4 AV nonstructural proteins, an AV RNA polymerase recognition sequence, and a 3' polyadenylate tract, where the nucleic acid sequence which operably encodes all 4 AV nsP's contains a mutation in at least one nsP that is a mutation in nsP1 residues 346, 441, 473, nsP2 residues 438, 622, 634, 715, nsP3 residues, 417, 456, 505, and nsP4 residue 266, as compared to wild-type; and

(14) an AV RNA vector replicon capable of translation in a eukaryotic system, comprising a 5' sequence which initiates transcription of AV RNA, a nucleic acid molecule which operably encodes all 4 AV nsP's, an AV RNA polymerase recognition sequence, and a 3' polyadenylate tract, where the nucleic acid sequence which operably encodes all 4 AV nsP's contains a mutation in at least one nsP that is a mutation in nsP1 residues 346, 441, 473, nsP2 residues 438, 622, 634, 715, nsP3 residues, 417, 456, 505, and nsP4 residue 266, as compared to wild-type.

ACTIVITY - Immunostimulatory; cytostatic; virucide; fungicide; antibacterial; antiparasitic. No suitable biological data is given.

MECHANISM OF ACTION - Vaccine.

USE - The AV's are used to infect dendritic cells, preferably human cells (claimed). A heterologous sequence can be introduced and expressed in human macrophages or antigen presenting cells in vivo and in vitro, for use in biological assays. The AV-based vector systems are used to generate an immune response to cancer or a pathogenic agent, such as, bacteria, fungi, parasites or viruses.

ADVANTAGE - The AV can be used to infect human dendritic cells, macrophages or antigen presenting cells that previously could not be infected using an AV or AV variant. The AV vectors are targeted directly to antigen presenting cells. Dwg.0/12

L11 ANSWER 9 OF 33 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER:

2000-452400 [39] WPIDS

CROSS REFERENCE:

2000-452401 [39]; 2000-465745 [39]

DOC. NO. CPI:

C2000-137949

TITLE:

Expression cassettes encoding the human immunodeficiency virus (HIV) Gag-containing polypeptide useful for vaccinating against HIV infections and acquired immunodeficiency syndrome

(AIDS).

DERWENT CLASS:

B04 C06 D16

INVENTOR (S):

BARNETT, S; GREER, C; HARTOG, K; LIAN, Y; LIU, H; SELBY, M; SRIVASTAVA, I; WALKER, C; ZUR MEGEDE, J

PATENT ASSIGNEE(S):

(CHIR) CHIRON CORP

COUNTRY COUNT:

89

PATENT INFORMATION:

PATENT NO KIND DATE WEEK LA PG

WO 2000039302 A2 20000706 (200039)* EN 390

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ TZ UG ZW

W: AE AL AM AT AU AZ BA BB BG BR BY CA CH CN CR CU CZ DE DK DM EE ES FI GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MA MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT TZ UA UG UZ VN YU ZA ZW

AU 2000022216 A 20000731 (200050)

APPLICATION DETAILS:

PATENT NO K	IND	API	LICATION	DATE
WO 2000039302	~ 		1999-US31245	
AU 2000033302				19991230

FILING DETAILS:

PATENT NO	KIND			PA	TENT NO
AU 20000222	16 A	Based	on	WO	200039302

PRIORITY APPLN. INFO: US 1999-168471 19991201; US 1998-114495 19981231

AN 2000-452400 [39] WPIDS

CR 2000-452401 [39]; 2000-465745 [39]

AB WO 200039302 A UPAB: 20001010

NOVELTY - Synthetic expression cassettes comprising nucleic acids encoding the human immunodeficiency virus (HIV) Gag-containing polypeptide, are new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are included for the following:

- (1) an expression cassette (I) comprising a polynucleotide sequence encoding a protein comprising a human immunodeficiency virus (HIV) Gag polypeptide (the polynucleotide sequence encoding the Gag polypeptide comprises a sequence with at least 90% sequence identity to a defined 60 nucleotide sequence (N1) given in the specification);
- (2) a **recombinant** expression system (II) for use in a host cell comprising (I) operably linked to control elements suitable or protein expression in the host;
 - (3) a cell (III) comprising (II);
- (4) a method (IV) for producing polypeptides including HIV Gag polypeptide sequences, comprising incubating (III) under conditions suitable for expression of the polypeptide;
- (5) a method (V) for producing virus-like particles (VLPs), comprising incubating (III) under conditions suitable for production

of VLPs; and

(6) a method (VI) for DNA vaccination of a subject, comprising introducing (II) into a subject under conditions suitable for gene expression.

ACTIVITY - Anti-HIV.

MECHANISM OF ACTION - Vaccine.

USE - The expression cassettes may be used for the recombinant expression of HIV Gag-polypeptides which may then be used to vaccinate against HIV infection and acquired immunodeficiency syndrome (AIDS).

Dwg.0/82

L11 ANSWER 10 OF 33 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER:

2000-431307 [37] WPIDS

DOC. NO. CPI:

C2000-131084

TITLE:

Novel recombinant vector useful as

immunogens and vaccines for stimulating and

enhancing immunological responses to target cells and antigens expresses multiple co-stimulatory

molecules such as B7-1, LFA-3, ICAM-1.

DERWENT CLASS:

B04 D16

INVENTOR(S):

HODGE, J; PANICALI, D; SCHLOM, J

PATENT ASSIGNEE(S):

(THER-N) THERION BIOLOGICS CORP; (USSH) US DEPT

HEALTH & HUMAN SERVICES

COUNTRY COUNT:

90

PATENT INFORMATION:

PATENT	NO	KIND	DATE	WEEK	LA	PG
					 _	

WO 2000034494 A1 20000615 (200037)* EN 188

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SL SZ TZ UG ZW

W:AEALAMATAUAZBABBBGBRBYCACHCNCRCUCZDEDKDMEEESFIGBGDGEGHGMHRHUIDILINISJPKEKGKPKRKZLCLKLRLSLTLULVMAMDMGMKMNMWMXNONZPLPTRORUSDSESGSISKSLTJTMTRTTTZUAUGUSUZVNYUZAZW

AU 2000016218 A 20000626 (200045)

APPLICATION DETAILS:

PATENT NO	KIND	AP	PLICATION	DATE
WO 2000034494	4 A1	WO	1999-US26866	19991112
ATT 200001621	8 Z	ΔIJ	2000-16218	19991112

FILING DETAILS:

PATENT NO KIND

PATENT NO

AM 2000016210 A Board on WO 200024

AU 2000016218 A Based on

WO 200034494

PRIORITY APPLN. INFO: US 1998-111582 19981209

AN 2000-431307 [37] WPIDS

AB WO 200034494 A UPAB: 20000807

NOVELTY - A recombinant vector (I), comprising foreign nucleic acid sequences encoding multiple co-stimulatory molecules (CM), or functional portions of them, is new.

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for the following:

- (1) a pharmaceutical composition, comprising (I) or a recombinant vector comprising a nucleic sequence encoding a target antigen or an immunological epitope, and a carrier;
 - (2) a host cell (II) infected, transfected or induced with (I);
- (3) a dendritic cell (III), or its precursor, comprising a foreign nucleic acid sequence encoding multiple CM;
- (4) a tumor cell (IV), or its precursor, comprising a foreign nucleic acid sequence encoding multiple CM;
- (5) a pharmaceutical composition, comprising (II) and optionally a exogenous source of target antigen or its immunological epitope;
- (6) a recombinant poxvirus (V) having integrated into its genome a foreign DNA encoding multiple CM which is produced by allowing a plasmid vector comprising the foreign DNA encoding multiple CM to undergo recombination with the parental poxvirus genome to produce recombinant poxvirus with the foreign DNA inserted into its genome;
- (7) a recombinant poxvirus (VI) having integrated into its genome a foreign DNA encoding leukocyte function-associated antigen (LFA)-3, intracellular adhesion molecule (ICAM)-1 and at least one B7 molecule, produced by allowing a vector comprising the DNA to undergo recombination with a parenteral poxvirus genome to produce the recombinant virus;
- (8) a pharmaceutical composition comprising (V), or (VI), and a carrier;
 - (9) a host cell (VII) infected with (V) or (VI);
- (10) a plasmid vector (VIII) comprising nucleic acid sequences encoding multiple CM or its functional portions;
- (11) a plasmid vector (IX) for recombination with a poxvirus designed to produce **recombinant** poxvirus capable of expressing foreign nucleic acid encoding LFA-3, ICAM-1, and at least one B7 molecule, comprising poxviral promoters, the DNA encoding the costimulatory molecules, and DNA sequences flanking the construct of the promoters and DNA, which are homologous to a region of the poxvirus genome into which the construct is to insert;
- (12) a kit for use in making (V) of (VI), comprising (VIII) or (IX), and optionally a parental poxvirus; and

(13) making (V), or (VI) which involves allowing (VIII), or (IX) to undergo recombination with a parental poxvirus genome to produce recombinant poxvirus with the foreign DNA inserted into its genome and a multiplicity of poxvirus promoters capable of controlling the expression of the foreign DNA.

ACTIVITY - Antitumor; Cytostatic; antibacterial; antiviral; antifungal; antiprotozoan; antiulcer; antiinflammatory; antiparasitic. The cytostatic activity of (I) was tested using mice. A four-gene vaccinia recombinant was constructed that contained the human CEA (carcinoembryonic antigen) gene and the B7-1, ICAM-1 (intercellular adhesion molecule-1) and LFA-3 (leukocyte function associated antigen-3) genes, designated rV-CEA/TRICOM. Six to eight-week-old female C57 BL/6 mice C57BL/6 mice transgenic for human CEA (Kass,E et al Cancer Res 59:676-683, 1999) were vaccinated by tail scarification with 107 plaque forming units rV-CEA, rV-CEA/B7-1 or rV-CEA/TRICOM, and spleens were harvested 22 days later. Lymphoproliferative activity of splenocytes was analyzed. Splenic T-cells of mice vaccinated with rV-TRICOM showed higher levels of CEA-specific stimulation compared with T-cells obtained from mice vaccinated with rV-CEA.

MECHANISM OF ACTION - Vaccine; gene therapy.

USE - (I) is useful for enhancing an cell-mediated or humoral immune response in an individual. (V) is used for enhancing an antigen specific T-cell response in an individual to a target antigen. (II) is used in immunotherapy for treating or preventing diseases caused by viruses, bacteria, protozoans, parasites, premalignant cells and tumor cells. (III) or (IV) are also used in enhancing an immune response in an individual. (I) is used for making a progenitor DC, or DC derived from bone marrow or peripheral blood mononuclear cells that overexpresses multiple CM which involves providing a cell with (I) for a sufficient period of time to cause of overexpression of multiple CM. (II) is also used for assessing the efficacy of a vaccine against target antigen. (II) is used for screening novel immunogenic peptides from a combinatorial peptide library source which involves, pulsing APC infected with (I) with the peptides to form peptide-pulsed APC, and measuring lymphoid immunoreactivity in the presence of peptide-pulsed APC. An enhanced immunoreactivity is indicative of an immunogenic peptide-pulsed APC. (All claimed). (I) is useful as an immunogen and vaccine against pathogenic microorganisms and cancer and as diagnostic agents. The recombinant vector can be used to treat or prevent preneoplastic or hyperplastic states such as colon polyps, Crohn's disease, ulcerative colitis and breast lesions.

ADVANTAGE - The enhancement of the immunological response using the **recombinant** vectors expressing multiple costimulatory molecules is synergistic compared to the use of a single costimulatory molecule, or the use of two costimulatory molecules in enhancing immunological responses. The magnitude of the immune

response against the target antigen, epitope, or cells expressing target antigen obtained using the **recombinant** vector is significantly greater than that achieved using systems employing a single or a double costimulatory molecule (claimed).

DESCRIPTION OF DRAWING(S) - The figure shows the genomic structure of pT5032 comprising nucleic acid sequences encoding murine LFA-3 (leukocyte function associated antigen-3), ICAM-1 (intercellular adhesion molecule-1), B7.1, flanked by portions of the Honda M region of the vaccinia genome.

Dwg.1/54

L11 ANSWER 11 OF 33 MEDLINE

ACCESSION NUMBER: 2001069343 MEDLINE

DOCUMENT NUMBER: 20519611 PubMed ID: 10924501

TITLE: The metabotropic GABAB receptor directly interacts

with the activating transcription factor 4.

AUTHOR: Nehring R B; Horikawa H P; El Far O; Kneussel M;

Brandstatter J H; Stamm S; Wischmeyer E; Betz H;

Karschin A

CORPORATE SOURCE: Department of Molecular Neurobiology of Signal

Transduction, Max Planck Institute for Biophysical

Chemistry, 37070 Gottingen, Germany.

SOURCE: JOURNAL OF BIOLOGICAL CHEMISTRY, (2000 Nov 10) 275

(45) 35185-91.

Journal code: HIV. ISSN: 0021-9258.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200101

ENTRY DATE: Entered STN: 20010322

Last Updated on STN: 20010322 Entered Medline: 20010104

G protein-coupled receptors regulate gene expression by cellular AB signaling cascades that target transcription factors and their recognition by specific DNA sequences. In the central nervous system, heteromeric metabotropic gamma-aminobutyric acid type B (GABA(B)) receptors through adenylyl cyclase regulate cAMP levels, which may control transcription factor binding to the cAMP response element. Using yeast-two hybrid screens of rat brain libraries, we now demonstrate that GABA(B) receptors are engaged in a direct and specific interaction with the activating transcription factor 4 (ATF-4), a member of the cAMP response element-binding protein /ATF family. As confirmed by pull-down assays, ATF-4 associates via its conserved basic leucine zipper domain with the C termini of both GABA(B) receptor (GABA(B)R) 1 and GABA(B)R2 at a site which serves to assemble these receptor subunits in heterodimeric complexes. Confocal fluorescence microscopy shows that GABA(B)R and ATF-4 are

strongly coclustered in the soma and at the dendritic membrane surface of both cultured hippocampal neurons as well as retinal amacrine cells in vivo. In oocyte coexpression assays short term signaling of GABA(B)Rs via G proteins was only marginally affected by the presence of the transcription factor, but ATF-4 was moderately stimulated in response to receptor activation in in vivo reporter assays. Thus, inhibitory metabotropic GABA(B)Rs may regulate activity-dependent gene expression via a direct interaction with ATF-4.

MEDLINE

DUPLICATE 2 L11 ANSWER 12 OF 33 MEDLINE

ACCESSION NUMBER: 2001083046

> 20541985 PubMed ID: 11090185

DOCUMENT NUMBER: Infection of human dendritic cells by a TITLE:

sindbis virus replicon vector is determined by a single amino acid substitution in the E2

glycoprotein.

Gardner J P; Frolov I; Perri S; Ji Y; MacKichan M L; **AUTHOR:**

zur Megede J; Chen M; Belli B A; Driver D A; Sherrill

S; Greer C E; Otten G R; Barnett S W; Liu M A;

Dubensky T W; Polo J M

Vaccines & Gene Therapy, Chiron Corporation, CORPORATE SOURCE:

Emeryville, California 94608, USA.

JOURNAL OF VIROLOGY, (2000 Dec) 74 (24) 11849-57. SOURCE:

Journal code: KCV. ISSN: 0022-538X.

United States PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

Priority Journals FILE SEGMENT:

200101 ENTRY MONTH:

Entered STN: 20010322 ENTRY DATE:

> Last Updated on STN: 20010322 Entered Medline: 20010111

The ability to target antigen-presenting cells with vectors encoding AB desired antigens holds the promise of potent prophylactic and therapeutic vaccines for infectious diseases and cancer. Toward this goal, we derived variants of the prototype alphavirus, Sindbis virus (SIN), with differential abilities to infect human dendritic cells. Cloning and sequencing of the SIN variant genomes revealed that the genetic determinant for human dendritic cell (DC) tropism mapped to a single amino acid substitution at residue 160 of the envelope glycoprotein E2. Packaging of SIN replicon vectors with the E2 glycoprotein from a DC -tropic variant conferred a similar ability to efficiently infect immature human DC, whereupon those DC were observed to undergo rapid activation and maturation. The SIN

replicon particles infected skin-resident mouse DC in

vivo, which subsequently migrated to the draining lymph nodes and upregulated cell surface expression of major histocompatibility complex and costimulatory molecules. Furthermore, SIN replicon particles encoding human immunodeficiency virus type 1 p55 (Gag) elicited robust Gag-specific T-cell responses in vitro and in vivo, demonstrating that infected DC maintained their ability to process and present replicon-encoded antigen. Interestingly, human and mouse DC were differentially infected by selected SIN variants, suggesting differences in receptor expression between human and murine DC. Taken together, these data illustrate the tremendous potential of using a directed approach in generating alphavirus vaccine vectors that target and activate antigen-presenting cells, resulting in robust antigen-specific immune responses.

L11 ANSWER 13 OF 33 MEDLINE DUPLICATE 3

ACCESSION NUMBER: 2000129590 MEDLINE

DOCUMENT NUMBER: 20129590 PubMed ID: 10666209

DOCUMENT NUMBER: 20129590 Publied 1D: 10060209

TITLE: Stimulation of cytotoxic T cells against idiotype

immunoglobulin of malignant lymphoma with protein-pulsed or idiotype-transduced

dendritic cells.

AUTHOR: Osterroth F; Garbe A; Fisch P; Veelken H

CORPORATE SOURCE: Departments of Hematology/Oncology and Pathology,

Freiburg University Medical Center, Freiburg,

Germany.

SOURCE: BLOOD, (2000 Feb 15) 95 (4) 1342-9.

Journal code: A8G; 7603509. ISSN: 0006-4971.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

ENTRY MONTH: 200003

ENTRY DATE: Entered STN: 20000327

Last Updated on STN: 20000327 Entered Medline: 20000314

AB Because of their hypervariable regions and somatic mutations, the antigen receptor molecules of lymphomas (idiotypes) are tumor-specific antigens and attractive targets for antilymphoma immunotherapy. For the optimal induction of human idiotype-specific cytotoxic T cells (CTL), idiotype was presented to CD8(+) peripheral blood mononuclear cells by monocyte-derived autologous dendritic cells (DC) after the endocytosis of idiotype protein or by idiotype-expressing DC.

Recombinant idiotype was obtained as a functionally folded Fab fragment by periplasmic expression in Escherichia coli. Idiotype-expressing DC were generated by transduction with recombinant Semliki forest virus vectors

encompassing heavy- or light-chain idiotype genes. Autologous lymphoblastoid cell lines stably transfected with Epstein-Barr virus-based idiotype expression vectors were used as target cells to detect idiotype-specific lysis. CTL stimulated with idiotype-loaded DC showed strong specific, CD8-mediated, and major histocompatibility complex (MHC) class I-restricted cytotoxicity against autologous heavy- and light-chain idiotype. In contrast, stimulation with idiotype-transduced DC resulted in only moderate natural killer cell activity. These data confirm the existence of idiotype-specific CTL in patients with lymphoma, define a "good manufacturing practice"-compatible protocol for the generation of these cells without the requirement of viable lymphoma cells, and favor the processing of exogenous antigen over DC transduction for the induction of MHC I-restricted CTL against idiotypes with unknown antigenicity. (Blood. 2000;95:1342-1349)

DUPLICATE 4 L11 ANSWER 14 OF 33 MEDLINE

ACCESSION NUMBER: 2000091347

MEDLINE PubMed ID: 10623754

20091347 DOCUMENT NUMBER:

TITLE:

Role of dendritic cell targeting in

Venezuelan equine encephalitis

virus pathogenesis.

MacDonald G H; Johnston R E **AUTHOR:**

Department of Microbiology and Immunology, University CORPORATE SOURCE:

of North Carolina at Chapel Hill School of Medicine,

Chapel Hill, North Carolina 27599-7290, USA...

gmacd@med.unc.edu

CONTRACT NUMBER: A122186 (NIAID)

F32-AI09778 (NINDS)

NS26681

JOURNAL OF VIROLOGY, (2000 Jan) 74 (2) 914-22. SOURCE:

Journal code: KCV; 0113724. ISSN: 0022-538X.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

English LANGUAGE:

FILE SEGMENT: Priority Journals

200002 ENTRY MONTH:

Entered STN: 20000218 ENTRY DATE:

> Last Updated on STN: 20000218 Entered Medline: 20000208

The initial steps of Venezuelan equine AB

encephalitis virus (VEE) spread from inoculation in the skin to the draining lymph node have been characterized. By using green fluorescent protein and immunocytochemistry, dendritic cells in the draining lymph node were determined to be the primary target of VEE infection in the first 48 h following inoculation. VEE viral replicon particles, which can undergo only one round of infection, identified Langerhans cells to be the initial set of

> Shears 308-4994 Searcher

cells infected by VEE directly following inoculation. These cells are resident dendritic cells in the skin, which migrate to the draining lymph node following activation. A point mutation in the E2 glycoprotein gene of VEE that renders the virus avirulent and compromises its ability to spread beyond the draining lymph blocked the appearance of virally infected dendritic cells in the lymph node in vivo. A second-site suppressor mutation that restores viral spread to lymphoid tissues and partially restore virulence likewise restored the ability of VEE to infect dendritic cells in vivo.

L11 ANSWER 15 OF 33 MEDLINE

ACCESSION NUMBER: 2001110759 MEDLINE

DOCUMENT NUMBER: 20565030 PubMed ID: 11112802

TITLE: Imaging high-resolution structure of GFP-expressing

neurons in neocortex in vivo.

AUTHOR: Chen B E; Lendvai B; Nimchinsky E A; Burbach B; Fox

K; Svoboda K

CORPORATE SOURCE: Howard Hughes Medical Institute, Cold Spring Harbor

Laboratory, Cold Spring Harbor, New York 11724, USA.

SOURCE: LEARNING AND MEMORY, (2000 Nov-Dec) 7 (6) 433-41.

Journal code: DAB. ISSN: 1072-0502.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200102

ENTRY DATE: Entered STN: 20010322

Last Updated on STN: 20010322 Entered Medline: 20010202

AB To detect subtle changes in neuronal morphology in response to changes in experience, one must image neurons at high resolution in vivo over time scales of minutes to days. We accomplished this by infecting postmitotic neurons in rat and mouse barrel cortex with a Sindbis virus carrying the gene for enhanced green

fluorescent protein. Visualized with 2-photon excitation laser scanning microscopy, infected neurons showed bright fluorescence that was distributed homogeneously throughout the cell, including axonal and dendritic arbors. Single dendritic

spines could routinely be resolved and their morphological dynamics visualized. Viral infection and imaging were achieved throughout postnatal development up to early adulthood (P 8-30), although the viral efficiency of infection decreased with age. This relatively noninvasive method for fluorescent labeling and imaging of neurons allows the study of morphological dynamics of neocortical neurons and their circuits in vivo.

L11 ANSWER 16 OF 33 MEDLINE

ACCESSION NUMBER: 2000454443 MEDLINE

DOCUMENT NUMBER: 20439484 PubMed ID: 10985351

TITLE: Mutagenesis reveals a role for ABP/GRIP binding to

GluR2 in synaptic surface accumulation of the AMPA

receptor.

AUTHOR: Osten P; Khatri L; Perez J L; Kohr G; Giese G; Daly

C; Schulz T W; Wensky A; Lee L M; Ziff E B

CORPORATE SOURCE: Max-Planck Institute for Medical Research, Department

of Molecular Neurobiology, Heidelberg, Germany...

posten@mpimf-heidelberg.mpg.de

CONTRACT NUMBER: AG13620 (NIA)

SOURCE: NEURON, (2000 Aug) 27 (2) 313-25.

Journal code: AN8; 8809320. ISSN: 0896-6273.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200009

ENTRY DATE: Entered STN: 20001005

Last Updated on STN: 20001005 Entered Medline: 20000926

AB We studied the role of PDZ proteins GRIP, ABP, and PICK1 in GluR2 AMPA receptor trafficking. An epitope-tagged MycGluR2 subunit, when expressed in hippocampal cultured neurons, was specifically targeted to the synaptic surface. With the mutant MycGluR2delta1-10, which lacks the PDZ binding site, the overall dendritic intracellular transport and the synaptic surface targeting were not affected. However, over time, Myc-GluR2delta1-10 accumulated at synapses significantly less than MycGluR2. Notably, a single residue substitution, S880A, which blocks binding to ABP/GRIP but not to PICK1, reduced synaptic accumulation to the same extent as the PDZ site truncation. We conclude that the association of GluR2 with ABP and/or GRIP but not PICK1 is essential for maintaining the synaptic surface accumulation of the receptor, possibly by limiting its endocytotic rate.

L11 ANSWER 17 OF 33 SCISEARCH COPYRIGHT 2001 ISI (R)

ACCESSION NUMBER: 2000:737596 SCISEARCH

THE GENUINE ARTICLE: 357CC

TITLE: Recombinant viruses as a tool for

therapeutic vaccination against human cancers

AUTHOR: Bonnet M C; Tartaglia J; Verdier F; Kourilsky P;

Lindberg A; Klein M; Moingeon P (Reprint)

CORPORATE SOURCE: AVENTIS PASTEUR, CAMPUS MERIEUX, 1541 AVE MARCEL

MERIEUX, F-69280 MARCY LETOILE, FRANCE (Reprint); AVENTIS PASTEUR, F-69280 MARCY LETOILE, FRANCE; AVENTIS PASTEUR, N YORK, ON, CANADA; INST PASTEUR,

PARIS, FRANCE

COUNTRY OF AUTHOR: FRANCE; CANADA

SOURCE: IMMUNOLOGY LETTERS, (15 SEP 2000) Vol. 74, No. 1,

Sp. iss. SI, pp. 11-25.

Publisher: ELSEVIER SCIENCE BV, PO BOX 211, 1000 AE

AMSTERDAM, NETHERLANDS.

ISSN: 0165-2478.

DOCUMENT TYPE: General Review; Journal

FILE SEGMENT: LIFE
LANGUAGE: English
REFERENCE COUNT: 105

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

Viral vectors can be used to express a variety of genes in vivo, AB that encode tumor associated antigens, cytokines, or accessory molecules. For vaccination pur poses, the ideal viral vector should be safe and enable efficient presentation of expressed antigens to the immune system. It should also exhibit low intrinsic immunogenicity to allow for its re-administration in order to boost relevant specific immune responses. Furthermore, the vector system must meet criteria that enable its industrialization. The characteristics of the most promising viral vectors, including retroviruses, poxviruses, adenoviruses, adeno-associated viruses, herpes simplex viruses, and alphaviruses, will be reviewed in this communication. Such recombinant viruses have been successfully used in animal models as therapeutic cancer vaccines. Based on these encouraging results, a series of clinical studies, reviewed herein, have been undertaken. Human clinical trials, have as of today, allowed investigators to establish that recombinant viruses can be safely used in cancer patients, and that such recombinants call break immune tolerance against tumor-associated antigens. These promising results are now leading to improved immunization protocols associating recombinant viruses with alternate antigen-presentation platforms (prime-boost regimens), in order to elicit broad tumor-specific immune responses (humoral and cellular) against multiple target antigens. (C) 2000 Elsevier Science B.V. All rights reserved.

L11 ANSWER 18 OF 33 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: 2001:88317 BIOSIS DOCUMENT NUMBER: PREV200100088317

TITLE: Viral gene transfer into neurons from hippocampal

slices: comparison of Semliki

Forest virus, adeno-associated virus, and

measles virus.

AUTHOR(S): Ehrengruber, M. U. (1); Hennou, S.; Lundstrom, K.;

Bueeler, H.; Naim, H. Y.; Gaehwiler, B. H.

CORPORATE SOURCE: (1) Univ Zurich, Zurich Switzerland

SOURCE: Society for Neuroscience Abstracts, (2000) Vol. 26,

No. 1-2, pp. Abstract No.-329.1. print.

Meeting Info.: 30th Annual Meeting of the Society of Neuroscience New Orleans, LA, USA November 04-09,

2000 Society for Neuroscience

. ISSN: 0190-5295.

DOCUMENT TYPE: Conference
LANGUAGE: English
CUMMARY LANGUAGE: English

SUMMARY LANGUAGE: English Viral vectors are useful to transfer cDNA into neurons. As diverse viruses have specific biological profiles, the ideal vector choice depends on both the gene and test system under study. We have previously shown that Semliki Forest virus (SFV) and Sindbis virus, two closely related positive-strand RNA viruses, efficiently infect neurons. In the present study, we characterized distinct vectors in rat hippocampal slice cultures. The following recombinant viruses encoding GFP were injected into stratum pyramidale: i) SFV, ii) adeno-associated virus (AAV), a single-strand DNA virus, and iii) measles virus (MV), a negative-strand RNA virus. All viruses efficiently infected pyramidal cells. GFP fluorescence was found in dendrites of up to the fourth order and in dendritic spines. SFV infected more pyramidal cells (apprx90% of all GFP-positive cells) than either AAV or MV (apprx65%). AAV-mediated GFP expression was more neuron-specific (apprx90%) when a PDGF rather than CMV promoter was used. GFP expression occurred rapidly but was transient for SFV (max. at 1-2 d post-infection, p.i.), increased slowly (from 5 d p.i.) but remained stable with AAV, and was fast (1-2 d p.i.) and persistent with MV. Resting membrane potentials and conductances as well as firing properties of pyramidal cells were normal at 2 and 28 d p.i. for SFV and AAV, respectively. We conclude that SFV is valuable for short-term, AAV for long-term, and MV for both shortand long-term gene transfer into pyramidal cells from hippocampal slices.

L11 ANSWER 19 OF 33 WPIDS COPYRIGHT 2001 DERWENT INFORMATION LTD

ACCESSION NUMBER: 1999-395093 [33] WPIDS

DOC. NO. CPI:

C1999-116132

TITLE:

Using new Venezuelan equine encephalitis virus vectors.

DERWENT CLASS:

B04 D16

INVENTOR(S):

HIPPENMEYER, P J

PATENT ASSIGNEE(S):

(SEAR) SEARLE & CO G D

COUNTRY COUNT:

85

PATENT INFORMATION:

RW: AT BE CH CY DE DK EA ES FI FR GB GH GM GR IE IT KE LS LU MC MW NL OA PT SD SE SZ UG ZW

W: AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI
GB GD GE GH GM HR HU ID IL IN IS JP KE KG KP KR KZ LC LK LR
LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI
SK SL TJ TM TR TT UA UG US UZ VN YU ZW

AU 9917106 A 19990705 (199948)

EP 1039926 A1 20001004 (200050) EN

R: AT BE CH DE DK ES FI FR GB GR IE IT LI LU NL PT SE

APPLICATION DETAILS:

PATENT NO	KIND	APPLICATION	DATE
WO 9930734	A1	WO 1998-US25725	19981214
AU 9917106	A	AU 1999-17106	19981214
EP 1039926	A1	EP 1998-961904	19981214
		WO 1998-US25725	19981214

FILING DETAILS:

PATENT	NO	KIND			PAT	ENT NO
AU 9917	7106	A	Based	on	WO	9930734
EP 1039	926	A1	Based	on	WO	9930734

PRIORITY APPLN. INFO: US 1997-68080 19971218

AN 1999-395093 [33] WPIDS

AB WO 9930734 A UPAB: 19990819

NOVELTY - Venezuelan equine encephalitis (VEE) virus vectors can be used to express tumor-associated antigens and cytokines, and thus induce immunity to cancer.

DETAILED DESCRIPTION - A method of protecting or treating a subject against primary or metastatic neoplastic diseases, is new and comprises administering an effective amount of recombinant VEE virus vector (I) which comprises at least one attenuating mutation, and a heterologous DNA segment comprising a promoter operably-linked to a DNA encoding a protein or peptide effective for treating the disease. INDEPENDENT CLAIMS are also included for:

- (1) a method of modulating tumours in a patient with (I);
- (2) a method of inhibiting proliferation of tumor cells with
 (I);
- (3) a process of inhibiting elevated levels of tumor cells, comprising administering to a host in need a therapeutically effective amount of (I) in unit dosage form;
- (4) a process of treating primary or metastatic neoplastic diseases administering to a mammalian host in need a therapeutically

effective amount of (I) in unit dosage form, alternatively, a tumor-associated peptide is administered;

- (5) a method of inhibiting the production of tumor cells in a patient comprising administering an effective amount of a tumor-associated peptide;
- (6) a DNA comprising a cDNA clone coding for an infectious VEE virus RNA transcript and a heterologous promoter positioned upstream from the cDNA clone and operably associated therewith, further comprising at least one attenuating mutation and containing the nucleotide sequence encoding a TAA, a TAA peptide, or a natural or synthetic cytokine operably linked to a promoter;
- (7) an infectious VEE RNA transcript encoded by the cDNA of (6);
- (8) a method of treating a subject against primary or metastatic neoplastic diseases by infecting a subjects dendritic cells with a the transcript of (7);
- (9) an inoculum comprising an effective amount of nucleic acid encoding the protein encoded by the cDNA of (6) dissolved or dispersed in an aqueous physiologically tolerable or pharmaceutically-acceptable diluent;
- (10) a pharmaceutical composition comprising a therapeutically effective amount of (I) in a mixture with a pharmaceutically acceptable carrier, optionally further comprising an adjunctive agent selected from chemotherapeutic or immunotherapeutic agents.
- USE The VEE virus vectors of the invention can be used prevent, treat, and protect against primary and metastatic neoplastic diseases, especially tumors such as lung cancer, breast cancer, ovarian cancer, prostate cancer, pancreatic cancer, gastric cancer, colon cancer, renal cancer, bladder cancer, melanoma, hepatoma, sarcoma and lymphoma (all claimed).

ADVANTAGE - Cancer is a leading cause of death. Conventional cancer treatments consist of chemotherapy, radiotherapy and surgery. The VEE virus vectors of the invention circumvents these conventional therapies, and instead uses the natural defense system of the body against the cancer cell. Dwg.0/0

L11 ANSWER 20 OF 33 MEDLINE DUPLICATE 5

ACCESSION NUMBER:

1999289594

MEDLINE

DOCUMENT NUMBER:

99289594 PubMed ID: 10359835

TITLE:

Recombinant Semliki

Forest virus and Sindbis virus

efficiently infect neurons in hippocampal slice

AUTHOR:

Ehrengruber M U; Lundstrom K; Schweitzer C; Heuss C;

Schlesinger S; Gahwiler B H

CORPORATE SOURCE:

Brain Research Institute, University of Zurich, Winterthurerstrasse 190, CH-8057 Zurich

Searcher :

Shears

308-4994

Switzerland.. ehrengru@hifo.unizh.ch

PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF SOURCE:

THE UNITED STATES OF AMERICA, (1999 Jun 8) 96 (12)

7041-6.

Journal code: PV3; 7505876. ISSN: 0027-8424.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

199907

ENTRY DATE:

Entered STN: 19990715

Last Updated on STN: 19990715 Entered Medline: 19990708

Gene transfer into nervous tissue is a powerful tool for the AB analysis of gene function. By using a rat hippocampal slice culture preparation, we show here that Semliki Forest virus (SFV) and Sindbis virus (SIN) vectors are useful for the effective infection of neurons. The stratum pyramidale and/or the granular cell layer were injected with recombinant virus encoding beta-galactosidase (LacZ) or green fluorescent protein (GFP). By using low concentrations of injected SFV-LacZ or SIN-LacZ, we detected LacZ staining of pyramidal cells, interneurons, and granule cells. About 60% of the infected cells showed clear neuronal morphology; thus, relatively few glial cells expressed the transgene. Expression of GFP from SFV and SIN vectors gave similar results, with an even higher percentage (>90%) of the GFP-positive cells identified as neurons. Infected pyramidal cells were readily recognized in living slices, displaying GFP fluorescence in dendrites of up to fourth order and in dendritic spines. They appeared morphologically normal and viable at 1-5 days postinfection. We conclude that both SFV and SIN vectors efficiently transfer genes into neurons in hippocampal slice cultures. In combination with the GFP reporter, SFV and SIN vectors will allow the physiological examination of identified neurons that have been modified by overexpression or suppression of a specific gene product.

L11 ANSWER 21 OF 33 SCISEARCH COPYRIGHT 2001 ISI (R)

ACCESSION NUMBER:

1999:767135 SCISEARCH

THE GENUINE ARTICLE: 242TE

TITLE:

Induction of P815 tumor immunity by

recombinant Semliki Forest virus expressing the P1A gene

AUTHOR:

Colmenero P; Liljestrom P; Jondal M (Reprint)

CORPORATE SOURCE:

KAROLINSKA INST, MICROBIOL & TUMORBIOL CTR, BOX 280, S-17177 STOCKHOLM, SWEDEN (Reprint); KAROLINSKA

INST, MICROBIOL & TUMORBIOL CTR, S-17177 STOCKHOLM,

SWEDEN; SWEDISH INST INFECT DIS CONTROL, DEPT

VACCINE RES, SOLNA, SWEDEN

COUNTRY OF AUTHOR:

SWEDEN

SOURCE:

GENE THERAPY, (OCT 1999) Vol. 6, No. 10, pp.

1728-1733.

Publisher: STOCKTON PRESS, HOUNDMILLS, BASINGSTOKE

RG21 6XS, HAMPSHIRE, ENGLAND.

ISSN: 0969-7128.

DOCUMENT TYPE:

Article; Journal

FILE SEGMENT:

LIFE

LANGUAGE:

English

REFERENCE COUNT:

54

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

The methylcholantrene-induced P815 mastocytoma tumor is derived from DBA/2 mice and expresses a weak tumor rejection antigen, P815A. The PIA gene, which encodes for the P815A antigen, is silent in most normal tissues with the exception of testis and placenta. These characteristics make P815 an interesting mouse model for the human MAGE-type tumor antigens. Recombinant Semliki

Forest virus particles (rSFV) were constructed that expressed variants of the P815 antigen. Such particles, when used for vaccination, express the antigen only transiently since the viral vector is incapable of productive replication. Nevertheless, mice vaccinated with rSFV generated strong CTL responses and were protected against P815 tumor challenge.

L11 ANSWER 22 OF 33 MEDLINE

ACCESSION NUMBER:

1999321252

MEDLINE

DOCUMENT NUMBER:

99321252 PubMed ID: 10395329

TITLE:
AUTHOR:

Cancer therapy using a self-replicating RNA vaccine. Ying H; Zaks T Z; Wang R F; Irvine K R; Kammula U S;

Marincola F M; Leitner W W; Restifo N P

CORPORATE SOURCE:

Surgery Branch, National Cancer Institute, Bethesda,

Maryland 20892-1502, USA.

SOURCE:

NATURE MEDICINE, (1999 Jul) 5 (7) 823-7.

Journal code: CG5; 9502015. ISSN: 1078-8956.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

199907

ENTRY DATE:

Entered STN: 19990806

Last Updated on STN: 19990806 Entered Medline: 19990729

'Naked' nucleic acid vaccines are potentially useful candidates for the treatment of patients with cancer, but their clinical efficacy has yet to be demonstrated. We sought to enhance the immunogenicity of a nucleic acid vaccine by making it 'self-replicating'. We accomplished this by using a gene encoding an RNA replicase

polyprotein derived from the Semliki forest virus, in combination with a model antigen. A single intramuscular injection of a self-replicating RNA immunogen elicited antigen-specific antibody and CD8+ T-cell responses at doses as low as 0.1 microg. Pre-immunization with a self-replicating RNA vector protected mice from tumor challenge, and therapeutic immunization prolonged the survival of mice with established tumors. The self-replicating RNA vectors did not mediate the production of substantially more model antigen than a conventional DNA vaccine did in vitro. However, the enhanced efficacy in vivo correlated with a caspase-dependent apoptotic death in transfected cells. This death facilitated the uptake of apoptotic cells by dendritic cells, providing a potential mechanism for enhanced immunogenicity. Naked, non-infectious, self-replicating RNA may be an excellent candidate for the development of new cancer vaccines.

L11 ANSWER 23 OF 33 EMBASE COPYRIGHT 2001 ELSEVIER SCI. B.V.DUPLICATE

ACCESSION NUMBER: 1999400725 EMBASE

TITLE: DNA and RNA-based vaccines: Principles, progress and

prospects.

AUTHOR: Leitner W.W.; Ying H.; Restifo N.P.

CORPORATE SOURCE: N.P. Restifo, National Cancer Institute, National

Institutes of Health, Building 10, Bethesda, MD

20892-1502, United States. restifo@nih.gov

SOURCE: Vaccine, (1999) 18/9-10 (765-777).

Refs: 142

ISSN: 0264-410X CODEN: VACCDE

PUBLISHER IDENT.: S 0264-410X(99)00271-6

COUNTRY: United Kingdom

DOCUMENT TYPE: Journal; General Review FILE SEGMENT: 022 Human Genetics

026 Immunology, Serology and Transplantation

029 Clinical Biochemistry

030 Pharmacology

037 Drug Literature Index

004 Microbiology

LANGUAGE: English
SUMMARY LANGUAGE: English

DNA vaccines were introduced less than a decade ago but have already been applied to a wide range of infectious and malignant diseases. Here we review the current understanding of the mechanisms underlying the activities of these new vaccines. We focus on recent strategies designed to enhance their function including the use of immunostimulatory (CpG) sequences, dendritic cells (DC), co-stimulatory molecules and cytokine- and chemokine-adjuvants. Although genetic vaccines have been significantly improved, they may not be sufficiently immunogenic for

the therapeutic vaccination of patients with infectious diseases or cancer in clinical trials. One promising approach aimed at dramatically increasing the immunogenicity of genetic vaccines involves making them 'self-replicating'. This can be accomplished by using a gene encoding RNA replicase, a polyprotein derived from alphaviruses, such as Sindbis virus.

Replicase-containing RNA vectors are significantly more immunogenic than conventional plasmids, immunizing mice at doses as low as 0.1 .mu.g of nucleic acid injected once intramuscularly. Cells transfected with 'self-replicating' vectors briefly produce large amounts of antigen before undergoing apoptotic death. This death is a likely result of requisite double-stranded (ds) RNA intermediates, which also have been shown to super-activate DC. Thus, the enhanced immunogenicity of 'self-replicating' genetic vaccines may be a result of the production of pro-inflammatory dsRNA, which mimics an RNA-virus infection of host cells. Copyright (C) 1999 Elsevier Science Ltd.

L11 ANSWER 24 OF 33 SCISEARCH COPYRIGHT 2001 ISI (R)

ACCESSION NUMBER: 1998:240119 SCISEARCH

THE GENUINE ARTICLE: ZC506

TITLE: Specific interactions between retrovirus Env and Gag

proteins in rat neurons

AUTHOR: Weclewicz K; Ekstrom M; Kristensson K; Garoff H

(Reprint)

CORPORATE SOURCE: KAROLINSKA INST, NOVUM, DEPT BIOSCI, S-14157

HUDDINGE, SWEDEN (Reprint); KAROLINSKA INST, NOVUM, DEPT BIOSCI, S-14157 HUDDINGE, SWEDEN; KAROLINSKA INST, DEPT NEUROSCI, S-17177 STOCKHOLM, SWEDEN

COUNTRY OF AUTHOR: SWEDEN

SOURCE: JOURNAL OF VIROLOGY, (APR 1998) Vol. 72, No. 4, pp.

2832-2845.

Publisher: AMER SOC MICROBIOLOGY, 1325 MASSACHUSETTS

AVENUE, NW, WASHINGTON, DC 20005-4171.

ISSN: 0022-538X. Article; Journal

DOCUMENT TYPE: A

LIFE English

LANGUAGE: En REFERENCE COUNT: 38

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

In this work we have studied the intracellular Localization properties of the Gag and Env proteins of Moloney murine leukemia virus (MLV) and human immunodeficiency virus (HIV) in dorsal root ganglion (DRG) neurons of rat, These neurons form thick bundles of axons, which facilitates protein localization studies by immunofluorescence analyses. When such neuron cultures were infected with recombinant Semliki Forest virus particles carrying the gag genes of either retrovirus, the expressed

Gag proteins were localized to both tile somatic and the axonal regions of the DRG neurons. In contrast, the Env proteins were confined only to the somatic region, When the Gag and Env proteins were coexpressed, the Gag proteins were also excluded from the axons. This effect of the Env proteins was shown to be dependent on the concentration of the Gag proteins in the neuron and also to be specific for homologous pairs of retrovirus proteins, Therefore, the results suggest that there are specific interactions between the Env and the Gag proteins of MLV and HIV in the DRG neurons.

L11 ANSWER 25 OF 33 SCISEARCH COPYRIGHT 2001 ISI (R)

ACCESSION NUMBER:

1998:794689 SCISEARCH

THE GENUINE ARTICLE: 126TN

TITLE:

Transfection of murine and human dendritic

cells with recombinant Semliki

Forest virus

AUTHOR:

Bungener L (Reprint); Schoemaker M; Wilschut J;

Daemen T

CORPORATE SOURCE:

UNIV GRONINGEN, DEPT PHYSIOL CHEM, GROINGEN UTRECHT

INST FRUG EXPLORAT, GRONINGEN, NETHERLANDS

COUNTRY OF AUTHOR:

NETHERLANDS

SOURCE:

JOURNAL OF LEUKOCYTE BIOLOGY, (OCT 1998) Supp. [2],

pp. K2-K2.

Publisher: FEDERATION AMER SOC EXP BIOL, 9650

ROCKVILLE PIKE, BETHESDA, MD 20814-3998.

ISSN: 0741-5400.

DOCUMENT TYPE:

Conference; Journal

FILE SEGMENT:

LIFE

LANGUAGE:

English

REFERENCE COUNT:

L11 ANSWER 26 OF 33 SCISEARCH COPYRIGHT 2001 ISI (R)

ACCESSION NUMBER:

97:879389 SCISEARCH

THE GENUINE ARTICLE: YG424

TITLE:

Transduction of human dendritic cells with

recombinant Semliki forest

virus coding for the ovarian carcinoma associated antigen alpha folate receptor for the generation of

cytotoxic T cells from cancer patients.

AUTHOR:

Albrecht B (Reprint); Kohler G; Mertelsmann R; Fisch

CORPORATE SOURCE:

UNIV FREIBURG, MED CTR, DEPT HEMATOL ONCOL, DEPT

PATHOL, D-7800 FREIBURG, GERMANY; UNIV TUBINGEN, DIV

IMMUNOL, TUBINGEN, GERMANY

COUNTRY OF AUTHOR:

GERMANY

SOURCE:

BLOOD, (15 NOV 1997) Vol. 90, No. 10, Part 1, Supp.

[1], pp. 2454-2454.

SAUNDERS CO, INDEPENDENCE SQUARE WEST

308-4994 Shears Searcher :

QP95768

CURTIS CENTER, STE 300, PHILADELPHIA, PA 19106-3399.

ISSN: 0006-4971.

DOCUMENT TYPE:

Conference; Journal

FILE SEGMENT:

LIFE; CLIN

LANGUAGE:

English

REFERENCE COUNT:

L11 ANSWER 27 OF 33 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER:

1998:68763 BIOSIS

PREV199800068763

DOCUMENT NUMBER: TITLE:

Transduction of human dendritic cells with

recombinant semliki forest

virus coding for the ovarian carcinoma associated antigen alpha folate receptor for the generation of

cytotoxic T cells from cancer patients.

AUTHOR (S):

Albrecht, B. (1); Koehler, G.; Mertelsmann, R.;

Fisch, P.

CORPORATE SOURCE:

(1) Univ. Freiburg Med. Cent., Dep. Hematol./Oncol.,

Freiburg Germany

SOURCE:

Blood, (Nov. 15, 1997) Vol. 90, No. 10 SUPPL. 1 PART

1, pp. 551A.

Meeting Info.: 39th Annual Meeting of the American Society of Hematology San Diego, California, USA December 5-9, 1997 The American Society of Hematology

. ISSN: 0006-4971.

DOCUMENT TYPE:

Conference

LANGUAGE:

English

L11 ANSWER 28 OF 33

MEDLINE

DUPLICATE 7

ACCESSION NUMBER:

96392394

DOCUMENT NUMBER:

96392394 PubMed ID: 8799182

MEDLINE

TITLE:

Expression and analysis of presentlin 1 in a human neuronal system: localization in cell bodies and

dendrites.

AUTHOR:

Cook D G; Sung J C; Golde T E; Felsenstein K M; Wojczyk B S; Tanzi R E; Trojanowski J Q; Lee V M;

Doms R W

CORPORATE SOURCE:

Department of Pathology and Laboratory Medicine, University of Pennsylvania, Philadelphia 19104, USA.

CONTRACT NUMBER:

P01 AG 11542 (NIA)

SOURCE:

PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE UNITED STATES OF AMERICA, (1996 Aug 20) 93 (17)

9223-8.

Journal code: PV3; 7505876. ISSN: 0027-8424.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH: 199610

ENTRY DATE: Entered STN: 19961219

Last Updated on STN: 19980206 Entered Medline: 19961031

AB Mutations in the recently identified presentlin 1 gene on chromosome 14 cause early onset familial Alzheimer disease (FAD). Herein we describe the expression and analysis of the protein coded by presentlin 1 (PS1) in NT2N neurons, a human neuronal model system.

PS1 was expressed using recombinant Semliki

Forest virions and detected by introduced antigenic tags or antisera to PS1-derived peptides. Immunoprecipitation revealed two major PS1 bands of approximately 43 and 50 kDa, neither of which were N-glycosylated or O-glycosylated. Immunoreactive PS1 was detected in cell bodies and dendrites of NT2N neurons but not in axons or on the cell surface. PS1 was also detected in BHK cells, where it was also intracellular and colocalized with calnexin, a marker for the rough endoplasmic reticulum. A mutant form of PS1 linked to FAD did not differ from the wild-type protein at the light microscopic level. The model system described here will enable studies of the function of PS1 in human neurons and the role of mutant PS1 in FAD.

L11 ANSWER 29 OF 33 MEDLINE

ACCESSION NUMBER: 97050828 MEDLINE

DOCUMENT NUMBER: 97050828 PubMed ID: 8895567

TITLE: The beta-amyloid domain is essential for axonal

sorting of amyloid precursor protein.

AUTHOR: Tienari P J; De Strooper B; Ikonen E; Simons M;

Weidemann A; Czech C; Hartmann T; Ida N; Multhaup G; Masters C L; Van Leuven F; Beyreuther K; Dotti C G

CORPORATE SOURCE: Cell Biology Programme, European Molecular Biology

Laboratories (EMBL), Heidelberg, Germany.

SOURCE: EMBO JOURNAL, (1996 Oct 1) 15 (19) 5218-29.

Journal code: EMB; 8208664. ISSN: 0261-4189.

PUB. COUNTRY: ENGLAND: United Kingdom

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199612

ENTRY DATE: Entered STN: 19970128

Last Updated on STN: 19980206 Entered Medline: 19961216

AB We have analysed the axonal sorting signals of amyloid precursor protein (APP). Wild-type and mutant versions of human APP were expressed in hippocampal neurons using the **Semliki**forest virus system. We show that wild-type APP and mutations implicated in Alzheimer's disease and another brain beta-amyloidosis are sorted to the axon. By analysis of deletion

mutants we found that the membrane-inserted APP ectodomain but not the cytoplasmic tail is required for axonal sorting. Systematic deletions of the APP ectodomain identified two regions required for axonal delivery: one encoded by exons 11-15 in the carbohydrate domain, the other encoded by exons 16-17 in the juxtamembraneous beta-amyloid domain. Treatment of the cells with the N-glycosylation inhibitor tunicamycin induced missorting of wild-type APP, supporting the importance of glycosylation in axonal sorting of APP. The data revealed a hierarchy of sorting signals on APP: the beta-amyloid-dependent membrane proximal signal was the major contributor to axonal sorting, while N-glycosylation had a weaker effect. Furthermore, recessive somatodendritic signals, most likely in the cytoplasmic tail, directed the protein to the dendrites when the ectodomain was deleted. Analysis of detergent solubility of APP and another axonally delivered protein, hemagglutinin, demonstrated that only hemagglutinin formed CHAPS-insoluble complexes, suggesting distinct mechanisms of axonal sorting for these two proteins. This study is the first delineation of sorting requirements of an axonally targeted protein in polarized neurons and indicates that the beta-amyloid domain plays a major role in axonal delivery of APP.

L11 ANSWER 30 OF 33 MEDLINE

ACCESSION NUMBER: 96032273 MEDLINE

DOCUMENT NUMBER: 96032273 PubMed ID: 7559765

TITLE: Intracellular routing of wild-type and mutated

polymeric immunoglobulin receptor in hippocampal

neurons in culture.

AUTHOR: de Hoop M; von Poser C; Lange C; Ikonen E; Hunziker

W; Dotti C G

CORPORATE SOURCE: European Molecular Biology Laboratory, Cell Biology

Program, Heidelberg, Germany.

SOURCE: JOURNAL OF CELL BIOLOGY, (1995 Sep) 130 (6) 1447-59.

Journal code: HMV; 0375356. ISSN: 0021-9525.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199511

ENTRY DATE: Entered STN: 19951227

Last Updated on STN: 19970203 Entered Medline: 19951106

AB Certain epithelial cells synthesize the polymeric immunoglobulin receptor (pIgR) to transport immunoglobulins (Igs) A and M into external secretions. In polarized epithelia, newly synthesized receptor is first delivered to the basolateral plasma membrane and is then, after binding the Ig, transcytosed to the apical plasma membrane, where the receptor-ligand complex is released by

proteolytic cleavage. In a previous work (Ikonen et al., 1993), we implied the existence of a dendro-axonal transcytotic pathway for the rabbit pIgR expressed in hippocampal neurons via the Semliki Forest Virus (SFV) expression system. By labeling surface-exposed pIgR in live neuronal cells, we now show (a) internalization of the receptor from the dendritic plasma membrane to the dendritic early endosomes, (b) redistribution of the receptor from the dendritic to the axonal domain, (c) inhibition of this movement by brefeldin A (BFA) and (d) stimulation by the activation of protein kinase C (PKC) via phorbol myristate acetate (PMA). In addition, we show that a mutant form of the receptor lacking the epithelial basolateral sorting signal is directly delivered to the axonal domain of hippocampal neurons. Although this mutant is internalized into early endosomes, no transcytosis to the dendrites could be observed. In epithelial Madin-Darby Canine Kidney (MDCK) cells, the mutant receptor could also be internalized into basolaterally derived early endosomes. These results suggest the existence of a dendro-axonal transcytotic pathway in neuronal cells which shares similarities with the basolateral to apical transcytosis in epithelial cells and constitute the basis for the future analysis of its physiological role.

L11 ANSWER 31 OF 33 MEDLINE DUPLICATE 8

ACCESSION NUMBER: 93367860 MEDLINE

DOCUMENT NUMBER: 93367860 PubMed ID: 8360951

mrmt n

TITLE: Expression of heterologous proteins in cultured rat

hippocampal neurons using the Semliki

Forest virus vector.

AUTHOR: Olkkonen V M; Liljestrom P; Garoff H; Simons K; Dotti

CG

CORPORATE SOURCE: Cell Biology Programme, European Molecular Biology

Laboratory, Heidelberg, Germany.

SOURCE: JOURNAL OF NEUROSCIENCE RESEARCH, (1993 Jul 1) 35 (4)

445-51.

Journal code: KAC; 7600111. ISSN: 0360-4012.

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199309

ENTRY DATE: Entered STN: 19931015

Last Updated on STN: 20000303 Entered Medline: 19930930

AB The Semliki Forest virus expression vector

(Liljestrom and Garoff: Bio/Technology 9:1356-1361, 1991) was tested in cultured rat hippocampal neurons using two Madin-Darby canine kidney (MDCK) cell membrane-associated proteins as reporters: rab8,

a small GTPase involved in post-Golgi vesicle transport, and VIP21, an integral membrane protein of caveolae, trans-Golgi network, and post-Golgi vesicles. Expression of the c-myc epitope-tagged proteins was visualized by immunofluorescence microscopy. The proteins were first detected in neurons after 3-4 hr infection by the recombinant viruses. The infection efficiency on neurons was high: after 6 hr infection at a multiplicity of one, 50-60% of the cells expressed the reporter proteins. The neurons tolerated the infection well up to 8 hr. Their polarized organization was not disturbed, as judged from morphology and from distribution of the dendritic MAP2 and axonal synaptophysin marker proteins. The Semliki Forest virus vector thus seems suitable for short-term expression of proteins in cultured neurons.

L11 ANSWER 32 OF 33 DUPLICATE 9 MEDLINE

94012966 MEDLINE ACCESSION NUMBER:

94012966 PubMed ID: 8408204 DOCUMENT NUMBER:

Protein transport to the dendritic plasma

TITLE:

membrane of cultured neurons is regulated by rab8p. Huber L A; de Hoop M J; Dupree P; Zerial M; Simons K; AUTHOR:

Dotti C

CORPORATE SOURCE: Cell Biology Programme, European Molecular Biology

Laboratory, Heidelberg, Germany.

JOURNAL OF CELL BIOLOGY, (1993 Oct) 123 (1) 47-55. SOURCE:

Journal code: HMV; 0375356. ISSN: 0021-9525.

United States PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

English LANGUAGE:

Priority Journals FILE SEGMENT:

199311 ENTRY MONTH:

Entered STN: 19940117 ENTRY DATE:

Last Updated on STN: 20000303

Entered Medline: 19931102

In the companion paper (Huber, L. A., S. W. Pimplikar, R. G. Parton, AB H. Virta, M. Zerial, and K. Simons. J. Cell Biol. 123:35-45) we reported that the small GTPase rab8p is involved in transport from the TGN to the basolateral plasma membrane in epithelia. In the present work we investigated the localization and function of rab8p in polarized hippocampal neurons. By immunofluorescence microscopy we found that rab8p localized preferentially in the somatodendritic domain, and was excluded from the axon. Double-labeling immunofluorescence showed that some of the rab8p co-localized in the dendrites with the Semliki Forest Virus glycoprotein **E2** (SFV-**E2**). An antisense oligonucleotide approach was used to investigate the role of rab8p in dendritic transport of newly synthesized viral glycoproteins. Antisense oligonucleotides corresponding to the initiation region of the rab8 coding sequence were added to the

cultured neurons for four days. This treatment resulted in a significant decrease in cellular levels of rab8p and transport of SFV-E2 from the cell body to the dendrites was significantly reduced. However, no effect was observed on axonal transport of influenza HA. From these results we conclude that rab8p is involved in transport of proteins to the dendritic surface in neurons.

L11 ANSWER 33 OF 33 CONFSCI COPYRIGHT 2001 CSA

1998:26500 CONFSCI ACCESSION NUMBER:

DOCUMENT NUMBER:

98-026500

TITLE:

Transduction of human dendritic cells with

recombinant Semliki Forest

virus coding for the ovarian carcinoma-associated antiqen [alpha] folate receptor for the generation of

cytotoxic T cells from cancer patients

AUTHOR: SOURCE:

Albrecht, B.; Koehler, G.; Mertelsmann, R.; Fisch, P. American Society for Hematology, 1200 19th Street, NW, Washington, DC 20036-2422; phone: 202-857-1118; fax: 202-857-1164; URL: http://www.hematology.org/, Abstracts available. Price \$40. Poster Paper No. 2454

Meeting Info.: 974 0032: 39th Annual Meeting of the American Society for Hematology (9740032). San Diego,

CA (USA). 5-9 Dec 1997. American Society for

Hematology.

DOCUMENT TYPE:

Conference

FILE SEGMENT:

DCCP English LANGUAGE:

FILE 'CAPLUS' ENTERED AT 11:47:36 ON 19 SEP 2001

658 S SFV(S)SEMLIKI OR SIN(S)SINDBIS OR VEE(S)VENEZUEL? L12

L13 17 S RRV(S)ROSS

12 S (L12 OR L13) AND (DENDRIT? OR DC(S)DENDRIT?) L14

L15 0 S L14 NOT L6

> J(FILE 'MEDLINE, BIOSIS, EMBASE, WPIDS, CONFSCI, SCISEARCH, JICST-EPLUS, JAPIÓ' ENTERED AT 11:50:14 ON 19 SEP 2001)

42 S L14 L16

L17 23 S L16 NOT L10

O S L17 AND ("E2" OR RESIDUE (5A) 160 OR RECOMBINAN?) L18

=> fil hom

FILE 'HOME' ENTERED AT 11:51:18 ON 19 SEP 2001

Searcher Shears

308-4994